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# 1 Trans-ancestry meta-analyses identify novel rare and common variants associated with blood

## 2 pressure and hypertension

3 Praveen Surendran <sup>1,167</sup>, Fotios Drenos <sup>2,3,167</sup>, Robin Young <sup>1,167</sup>, Helen Warren <sup>4,5,167</sup>, James P Cook <sup>6,7,167</sup>,  
4 Alisa K Manning <sup>8,9,10,167</sup>, Niels Grarup <sup>11,167</sup>, Xueling Sim <sup>12,13,14,167</sup>, Daniel R Barnes <sup>1</sup>, Kate Witkowska <sup>4,5</sup>,  
5 James R Staley <sup>1</sup>, Vinicius Tragante <sup>15</sup>, Taru Tukiainen <sup>8,9,16</sup>, Hanieh Yaghootkar <sup>17</sup>, Nicholas Masca <sup>18,19</sup>,  
6 Daniel F Freitag <sup>1</sup>, Teresa Ferreira <sup>20</sup>, Olga Giannakopoulou <sup>21</sup>, Andrew Tinker <sup>21,5</sup>, Magdalena Harakalova  
7 <sup>15</sup>, Evelin Mihailov <sup>22</sup>, Chunyu Liu <sup>23</sup>, Aldi T Kraja <sup>24,25</sup>, Sune Fallgaard Nielsen <sup>26</sup>, Asif Rasheed <sup>27</sup>, Maria  
8 Samuel <sup>27</sup>, Wei Zhao <sup>28</sup>, Lori L Bonnycastle <sup>29</sup>, Anne U Jackson <sup>13,12</sup>, Narisu Narisu <sup>29</sup>, Amy J Swift <sup>29</sup>,  
9 Lorraine Southam <sup>30,20</sup>, Jonathan Marten <sup>31</sup>, Jeroen R Huyghe <sup>13,12</sup>, Alena Stančáková <sup>32</sup>, Cristiano Fava <sup>33,34</sup>,  
10 Therese Ohlsson <sup>33</sup>, Angela Matchan <sup>30</sup>, Kathleen E Stirrups <sup>21,35</sup>, Jette Bork-Jensen <sup>11</sup>, Anette P Gjessing <sup>11</sup>,  
11 Jukka Kontto <sup>36</sup>, Markus Perola <sup>36,37,22</sup>, Susan Shaw-Hawkins <sup>4</sup>, Aki S Havulinna <sup>36</sup>, He Zhang <sup>38</sup>, Louise A  
12 Donnelly <sup>39</sup>, Christopher J Groves <sup>40</sup>, N William Rayner <sup>40,20,30</sup>, Matt J Neville <sup>40,41</sup>, Neil R Robertson <sup>20,40</sup>,  
13 Andrianos M Yiorkas <sup>42,43</sup>, Karl-Heinz Herzig <sup>44,45</sup>, Eero Kajantie <sup>36,46,47</sup>, Weihua Zhang <sup>48,49</sup>, Sara M  
14 Willems <sup>50</sup>, Lars Lannfelt <sup>51</sup>, Giovanni Malerba <sup>52</sup>, Nicole Soranzo <sup>53,35,54</sup>, Elisabetta Trabetti <sup>52</sup>, Niek  
15 Verweij <sup>55,9,56</sup>, Evangelos Evangelou <sup>48,57</sup>, Alireza Moayyeri <sup>48,58</sup>, Anne-Claire Vergnaud <sup>48</sup>, Christopher P  
16 Nelson <sup>18,19</sup>, Alaitz Poveda <sup>59,60</sup>, Tibor V Varga <sup>59</sup>, Muriel Caslake <sup>61</sup>, Anton JM de Craen <sup>62,63</sup>, Stella  
17 Trompet <sup>62,64</sup>, Jian'an Luan <sup>50</sup>, Robert A Scott <sup>50</sup>, Sarah E Harris <sup>65,66</sup>, David CM Liewald <sup>65,67</sup>, Riccardo  
18 Marioni <sup>65,66,68</sup>, Cristina Menni <sup>69</sup>, Aliko-Eleni Farmaki <sup>70</sup>, Göran Hallmans <sup>71</sup>, Frida Renström <sup>59,71</sup>, Jennifer  
19 E Huffman <sup>31,23</sup>, Maija Hassinen <sup>72</sup>, Stephen Burgess <sup>1</sup>, Ramachandran S Vasan <sup>23,73,74</sup>, Janine F Felix <sup>75</sup>,  
20 CHARGE-Heart Failure Consortium <sup>76</sup>, Maria Uria-Nickelsen <sup>77</sup>, Anders Malarstig <sup>78</sup>, Dermot F Reilly <sup>79</sup>,  
21 Maarten Hoek <sup>80</sup>, Thomas Vogt <sup>80,81</sup>, Honghuang Lin <sup>23,82</sup>, Wolfgang Lieb <sup>83</sup>, EchoGen Consortium <sup>76</sup>,  
22 Matthew Traylor <sup>84</sup>, Hugh F Markus <sup>84</sup>, METASTROKE Consortium <sup>76</sup>, Heather M Highland <sup>85</sup>, Anne E  
23 Justice <sup>85</sup>, Eirini Marouli <sup>21</sup>, GIANT Consortium <sup>76</sup>, Jaana Lindström <sup>36</sup>, Matti Uusitupa <sup>86,87</sup>, Pirjo  
24 Komulainen <sup>72</sup>, Timo A Lakka <sup>72,88,89</sup>, Rainer Rauramaa <sup>72,89</sup>, Ozren Polasek <sup>90,91</sup>, Igor Rudan <sup>90</sup>, Olov  
25 Rolandsson <sup>92</sup>, Paul W Franks <sup>59,92,93</sup>, George Dedoussis <sup>70</sup>, Timothy D Spector <sup>69</sup>, EPIC-InterAct  
26 Consortium <sup>76</sup>, Pekka Jousilahti <sup>36</sup>, Satu Männistö <sup>36</sup>, Ian J Deary <sup>65,67</sup>, John M Starr <sup>65,94</sup>, Claudia  
27 Langenberg <sup>50</sup>, Nick J Wareham <sup>50</sup>, Morris J Brown <sup>4</sup>, Anna F Dominiczak <sup>95</sup>, John M Connell <sup>39</sup>, J Wouter  
28 Jukema <sup>64,96</sup>, Naveed Sattar <sup>95</sup>, Ian Ford <sup>61</sup>, Chris J Packard <sup>61</sup>, Tõnu Esko <sup>22,97,8,9</sup>, Reedik Mägi <sup>22</sup>, Andres  
29 Metspalu <sup>22,98</sup>, Rudolf A de Boer <sup>99</sup>, Peter van der Meer <sup>99</sup>, Pim van der Harst <sup>99,100,101</sup>, Lifelines Cohort  
30 Study <sup>76</sup>, Giovanni Gambaro <sup>102</sup>, Erik Ingelsson <sup>103,104</sup>, Lars Lind <sup>103</sup>, Paul IW de Bakker <sup>105,106</sup>, Mattijs E  
31 Numans <sup>107,106</sup>, Ivan Brandslund <sup>108,109</sup>, Cramer Christensen <sup>110</sup>, Eva RB Petersen <sup>111</sup>, Eeva Korpi-Hyövälti  
32 <sup>112</sup>, Heikki Oksa <sup>113</sup>, John C Chambers <sup>48,49,114</sup>, Jaspal S Kooner <sup>49,115,114</sup>, Alexandra IF Blakemore <sup>42,43</sup>, Steve  
33 Franks <sup>116</sup>, Marjo-Riitta Jarvelin <sup>117,118,119,120</sup>, Lise L Husemoen <sup>121</sup>, Allan Linneberg <sup>121,122,123</sup>, Tea Skaaby  
34 <sup>121</sup>, Betina Thuesen <sup>121</sup>, Fredrik Karpe <sup>40,41</sup>, Jaakko Tuomilehto <sup>36,124,125,126</sup>, Alex SF Doney <sup>39</sup>, Andrew D  
35 Morris <sup>127</sup>, Colin NA Palmer <sup>39</sup>, Oddgeir Lingaas Holmen <sup>128,129</sup>, Kristian Hveem <sup>128,130</sup>, Cristen J Willer  
36 <sup>38,131,132</sup>, Tiinamaija Tuomi <sup>133,134</sup>, Leif Groop <sup>135,134</sup>, AnneMari Käräjämäki <sup>136,137</sup>, Aarno Palotie <sup>16,9,134,138</sup>,  
37 Samuli Ripatti <sup>134,139,30</sup>, Veikko Salomaa <sup>36</sup>, Dewan S Alam <sup>140</sup>, Abdulla al Shafi Majumder <sup>141</sup>, Emanuele Di  
38 Angelantonio <sup>1,54</sup>, Rajiv Chowdhury <sup>1</sup>, Mark I McCarthy <sup>40,41,20</sup>, Neil Poulter <sup>142</sup>, Alice V Stanton <sup>143</sup>, Peter  
39 Sever <sup>142</sup>, Philippe Amouyel <sup>144,145,146,147</sup>, Dominique Arveiler <sup>148</sup>, Stefan Blankenberg <sup>149,150</sup>, Jean Ferrières  
40 <sup>151</sup>, Frank Kee <sup>152</sup>, Kari Kuulasmaa <sup>36</sup>, Martina Müller-Nurasyid <sup>153,154,155</sup>, Giovanni Veronesi <sup>156</sup>, Jarmo  
41 Virtamo <sup>36</sup>, Panos Deloukas <sup>21,157</sup>, Wellcome Trust Case Control Consortium <sup>76</sup>, Paul Elliott <sup>117</sup>,  
42 Understanding Society Scientific Group <sup>76</sup>, Eleftheria Zeggini <sup>30</sup>, Sekar Kathiresan <sup>56,158,159,9</sup>, Olle Melander  
43 <sup>33</sup>, Johanna Kuusisto <sup>32</sup>, Markku Laakso <sup>32</sup>, Sandosh Padmanabhan <sup>95</sup>, David Porteous <sup>66</sup>, Caroline Hayward  
44 <sup>31</sup>, Generation Scotland <sup>160</sup>, Francis S Collins <sup>29</sup>, Karen L Mohlke <sup>161</sup>, Torben Hansen <sup>11</sup>, Oluf Pedersen <sup>11</sup>,  
45 Michael Boehnke <sup>13,12</sup>, Heather M Stringham <sup>13,12</sup>, EPIC-CVD Consortium <sup>76</sup>, Philippe Frossard <sup>27</sup>,  
46 Christopher Newton-Cheh <sup>56,158</sup>, CHARGE+ Exome Chip Blood Pressure Consortium <sup>76</sup>, Martin D Tobin <sup>6</sup>,

47 Børge Grønne Nordestgaard <sup>26</sup>, T2D-GENES Consortium <sup>76</sup>, GoT2DGenes Consortium <sup>76</sup>, ExomeBP  
48 Consortium <sup>76</sup>, CHD Exome+ Consortium <sup>76</sup>, Mark J Caulfield <sup>4,5</sup>, Anubha Mahajan <sup>20</sup>, Andrew P Morris  
49 <sup>20,7</sup>, Maciej Tomaszewski <sup>18,19,162</sup>, Nilesh J Samani <sup>18,19</sup>, Danish Saleheen <sup>28,27,1,167</sup>, Folkert W Asselbergs  
50 <sup>15,101,163,167</sup>, Cecilia M Lindgren <sup>164,9,20,167</sup>, John Danesh <sup>1,165,54,167</sup>, Louise V Wain <sup>6,167</sup>, Adam S Butterworth  
51 <sup>1,166,167</sup>, Joanna MM Howson <sup>1,167,168</sup>, Patricia B Munroe <sup>4,5,167,168</sup>

52

- 53 1. Cardiovascular Epidemiology Unit, Department of Public Health and Primary Care, University of  
54 Cambridge, Cambridge, UK
- 55 2. Medical Research Council Integrative Epidemiology Unit, School of Social and Community Medicine,  
56 University of Bristol, Oakfield House, Oakfield Grove, Bristol, UK
- 57 3. Centre for Cardiovascular Genetics, Institute of Cardiovascular Science, Rayne Building University  
58 College London, London, UK
- 59 4. Clinical Pharmacology, William Harvey Research Institute, Queen Mary University of London, London,  
60 UK
- 61 5. National Institute for Health Research Barts Cardiovascular Biomedical Research Unit, Queen Mary  
62 University of London, London, UK
- 63 6. Department of Health Sciences, University of Leicester, Leicester, UK
- 64 7. Department of Biostatistics, University of Liverpool, Liverpool, UK
- 65 8. Department of Genetics, Harvard Medical School, Boston, Massachusetts, USA
- 66 9. Program in Medical and Population Genetics, Broad Institute, 7 Cambridge Center, Cambridge,  
67 Massachusetts, USA
- 68 10. Department of Molecular Biology, Massachusetts General Hospital, Boston, Massachusetts, USA
- 69 11. The Novo Nordisk Foundation Center for Basic Metabolic Research, Faculty of Health and Medical  
70 Sciences, University of Copenhagen, Copenhagen, Denmark
- 71 12. Center for Statistical Genetics, University of Michigan, Ann Arbor, Michigan, USA
- 72 13. Department of Biostatistics, University of Michigan, Ann Arbor, Michigan, USA
- 73 14. Saw Swee Hock School of Public Health, National University of Singapore, National University Health  
74 System, Singapore
- 75 15. Department of Cardiology, University Medical Center Utrecht, Utrecht, The Netherlands
- 76 16. Analytic and Translational Genetics Unit, Department of Medicine, Massachusetts General Hospital,  
77 Boston, Massachusetts, USA
- 78 17. Genetics of Complex Traits, Institute of Biomedical and Clinical Science, University of Exeter Medical  
79 School, Exeter, UK
- 80 18. Department of Cardiovascular Sciences, University of Leicester, Leicester, UK
- 81 19. National Institute for Health Research Leicester Biomedical Research Unit in Cardiovascular Disease,  
82 Leicester, UK
- 83 20. Wellcome Trust Centre for Human Genetics, Nuffield Department of Medicine, University of Oxford,  
84 Oxford, UK
- 85 21. Heart Centre, William Harvey Research Institute, Barts and The London School of Medicine and  
86 Dentistry, Queen Mary University of London, London, UK
- 87 22. Estonian Genome Center, University of Tartu, Tartu, Estonia
- 88 23. National Heart, Lung, and Blood Institute's and Boston University's Framingham Heart Study,  
89 Framingham, Massachusetts, USA
- 90 24. Division of Statistical Genomics, Center for Genome Sciences and Systems Biology, Washington  
91 University School of Medicine, St. Louis, Missouri, USA
- 92 25. Department of Genetics, Washington University School of Medicine, St. Louis, Missouri, USA
- 93 26. Department of Clinical Biochemistry Herlev Hospital, Copenhagen University Hospital, Herlev,

94 Denmark  
95 27. Centre for Non-Communicable Diseases, Karachi, Pakistan  
96 28. Department of Biostatistics and Epidemiology, Perelman School of Medicine, University of  
97 Pennsylvania, Philadelphia, Pennsylvania, USA  
98 29. Medical Genomics and Metabolic Genetics Branch, National Human Genome Research Institute, NIH,  
99 Bethesda, Maryland, USA  
100 30. Wellcome Trust Sanger Institute, Genome Campus, Hinxton, UK  
101 31. Medical Research Council Human Genetics Unit, Medical Research Council Institute of Genetics and  
102 Molecular Medicine, University of Edinburgh, Edinburgh, UK  
103 32. Department of Medicine, University of Eastern Finland and Kuopio University Hospital, Kuopio,  
104 Finland  
105 33. University of Lund, Department of Clinical Sciences, Malmö, Sweden  
106 34. University of Verona, Department of Medicine, Verona, Italy  
107 35. Department of Haematology, University of Cambridge, Cambridge, UK  
108 36. Department of Health, National Institute for Health and Welfare, Helsinki, Finland  
109 37. Institute of Molecular Medicine FIMM, University of Helsinki, Finland  
110 38. Department of Internal Medicine, Division of Cardiovascular Medicine, University of Michigan, Ann  
111 Arbor, Michigan, USA  
112 39. Medical Research Institute, University of Dundee, Ninewells Hospital and Medical School, Dundee, UK  
113 40. Oxford Centre for Diabetes, Endocrinology, and Metabolism, Radcliffe Department of Medicine,  
114 University of Oxford, Oxford, UK  
115 41. National Institute for Health Research Oxford Biomedical Research Centre, Oxford University Hospital  
116 Trusts, Oxford, UK  
117 42. Section of Investigative Medicine, Imperial College London, London, UK  
118 43. Department of Life Sciences, Brunel University London, London, UK  
119 44. Institute of Biomedicine, Biocenter Oulu, University of Oulu, Oulu, Finland  
120 45. Department of Gastroenterology and Metabolism, Poznan University of Medical Sciences, Poznan,  
121 Poland  
122 46. Hospital for Children and Adolescents, Helsinki University Central Hospital and University of Helsinki,  
123 Helsinki, Finland  
124 47. Department of Obstetrics and Gynaecology, Oulu University Hospital and University of Oulu, Oulu,  
125 Finland  
126 48. Department of Epidemiology and Biostatistics, School of Public Health, Imperial College London,  
127 London, UK  
128 49. Department of Cardiology, Ealing Hospital, Middlesex, UK  
129 50. Medical Research Council Epidemiology Unit, University of Cambridge School of Clinical Medicine,  
130 Box 285 Institute of Metabolic Science, Cambridge Biomedical Campus, Cambridge, UK  
131 51. Department of Public Health and Caring Sciences, Uppsala University, Uppsala, Sweden  
132 52. Section of Biology and Genetics, Department of Neurosciences, Biomedicine and Movement Sciences,  
133 University of Verona, Verona, Italy  
134 53. Human Genetics, Wellcome Trust Sanger Institute, Hinxton, UK  
135 54. The National Institute for Health Research Blood and Transplant Research Unit in Donor Health and  
136 Genomics, University of Cambridge, Cambridge, UK  
137 55. University Medical Center Groningen, University of Groningen, Department of Cardiology, The  
138 Netherlands  
139 56. Center for Human Genetic Research, Massachusetts General Hospital, Boston, Massachusetts, USA  
140 57. Department of Hygiene and Epidemiology, University of Ioannina Medical School, Ioannina, Greece  
141 58. Farr Institute of Health Informatics Research, Institute of Health Informatics, University College

142 London, London, UK

143 59. Department of Clinical Sciences, Genetic and Molecular Epidemiology Unit, Lund University, Malmö,  
144 Sweden

145 60. Department of Genetics, Physical Anthropology and Animal Physiology, Faculty of Science and  
146 Technology, University of the Basque Country (UPV/EHU), Bilbao, Spain

147 61. University of Glasgow, Glasgow, UK

148 62. Department of Gerontology and Geriatrics, Leiden University Medical Center, Leiden, The Netherlands

149 63. Mr. De Craen suddenly passed away January 2016

150 64. Department of Cardiology, Leiden University Medical Center, Leiden, The Netherlands

151 65. Centre for Cognitive Ageing and Cognitive Epidemiology, University of Edinburgh, Edinburgh, UK

152 66. Centre for Genomic and Experimental Medicine, Medical Research Council Institute of Genetics and  
153 Molecular Medicine, University of Edinburgh, Edinburgh, UK

154 67. Department of Psychology, University of Edinburgh, Edinburgh, UK

155 68. Queensland Brain Institute, The University of Queensland, Brisbane, Queensland, Australia

156 69. Department of Twin Research and Genetic Epidemiology, King's College London, UK

157 70. Department of Nutrition and Dietetics, School of Health Science and Education, Harokopio University,  
158 Athens, Greece

159 71. Department of Biobank Research, Umeå University, Umeå, Sweden

160 72. Kuopio Research Institute of Exercise Medicine, Kuopio, Finland

161 73. Section of Cardiology, Department of Medicine, Boston University Schools of Medicine and Public  
162 Health, Boston, Massachusetts, USA

163 74. Sections of Preventive Medicine and Epidemiology, Department of Medicine, Boston University  
164 Schools of Medicine and Public Health, Boston, Massachusetts, USA

165 75. Department of Epidemiology, Erasmus MC, University Medical Center Rotterdam, Rotterdam, The  
166 Netherlands

167 76. A full list of members and affiliations appears in the Supplementary Note

168 77. Development Management and Planning, Pfizer Worldwide Research and Development

169 78. Pfizer Worldwide Research and Development, Stockholm, Sweden

170 79. Merck Research Laboratories, Genetics and Pharmacogenomics, Boston, Massachusetts, USA.

171 80. Merck Research Laboratories, Cardiometabolic Disease, Kenilworth, New Jersey, USA

172 81. CHDI Management/CHDI Foundation, Princeton, New Jersey, USA

173 82. Section of Computational Biomedicine, Department of Medicine, Boston University School of  
174 Medicine, Boston, Massachusetts, USA

175 83. Institute of Epidemiology and Biobank Popgen, Kiel University, Kiel, Germany

176 84. Neurology Unit, University of Cambridge, Cambridge Biomedical Campus, Cambridge, UK

177 85. University of North Carolina at Chapel Hill, Department of Epidemiology, Chapel Hill, North Carolina,  
178 USA

179 86. Department of Public Health and Clinical Nutrition, University of Eastern Finland, Finland

180 87. Research Unit, Kuopio University Hospital, Kuopio, Finland

181 88. Institute of Biomedicine/Physiology, University of Eastern Finland, Kuopio Campus, Finland

182 89. Department of Clinical Physiology and Nuclear Medicine, Kuopio University Hospital, Kuopio, Finland

183 90. Centre for Global Health Research, Usher Institute for Population Health Sciences and Informatics,  
184 University of Edinburgh, Edinburgh, UK

185 91. Faculty of Medicine, University of Split, Croatia

186 92. Department of Public Health and Clinical Medicine, Umeå University, Umeå, Sweden

187 93. Department of Nutrition, Harvard School of Public Health, Boston, Massachusetts, USA

188 94. Alzheimer Scotland Research Centre, University of Edinburgh, Edinburgh, UK

189 95. Institute of Cardiovascular and Medical Sciences, College of Medical, Veterinary and Life Sciences,

190 University of Glasgow, Glasgow, UK  
191 96. The Interuniversity Cardiology Institute of the Netherlands, Utrecht, The Netherlands  
192 97. Division of Endocrinology, Boston Children's Hospital, Boston, Massachusetts, USA  
193 98. Institute of Molecular and Cell Biology, Tartu, Estonia  
194 99. Department of Cardiology, University of Groningen, University Medical Center Groningen, Groningen,  
195 The Netherlands  
196 100. Department of Genetics, University of Groningen, University Medical Center Groningen, Groningen,  
197 The Netherlands  
198 101. Durrer Center for Cardiogenetic Research, ICIN-Netherlands Heart Institute, Utrecht, The Netherlands  
199 102. Division of Nephrology, Department of Internal Medicine and Medical Specialties, Columbus -  
200 Gemelli University Hospital, Catholic University, Rome, Italy  
201 103. Department of Medical Sciences, Molecular Epidemiology and Science for Life Laboratory, Uppsala  
202 University, Uppsala, Sweden  
203 104. Department of Medicine, Division of Cardiovascular Medicine, Stanford University School of  
204 Medicine, Stanford, California, USA  
205 105. Department of Medical Genetics, Center for Molecular Medicine, University Medical Center Utrecht,  
206 Utrecht, The Netherlands  
207 106. Department of Epidemiology, Julius Center for Health Sciences and Primary Care, University Medical  
208 Center Utrecht, Utrecht, The Netherlands  
209 107. Department of Public Health and Primary Care, Leiden University Medical Center, Leiden, The  
210 Netherlands  
211 108. Department of Clinical Biochemistry, Lillebaelt Hospital, Vejle, Denmark  
212 109. Institute of Regional Health Research, University of Southern Denmark, Odense, Denmark  
213 110. Medical Department, Lillebaelt Hospital, Vejle, Denmark  
214 111. Department of Clinical Immunology and Biochemistry, Lillebaelt Hospital, Vejle, Denmark  
215 112. South Ostrobothnia Central Hospital, Seinäjoki, Finland  
216 113. Tampere University Hospital, Tampere, Finland  
217 114. Imperial College Healthcare NHS Trust, London, UK  
218 115. National Heart and Lung Institute, Imperial College London, London, UK  
219 116. Institute of Reproductive and Developmental Biology, Imperial College London, London, UK  
220 117. Department of Epidemiology and Biostatistics, Medical Research Council Public Health England  
221 Centre for Environment and Health, School of Public Health, Faculty of Medicine, Imperial College  
222 London, St. Mary's Campus, London, UK  
223 118. Centre for Life Course Epidemiology, Faculty of Medicine, University of Oulu, Oulu, Finland  
224 119. Biocenter Oulu, University of Oulu, Oulu, Finland  
225 120. Unit of Primary Care, Oulu University Hospital, Oulu, Finland  
226 121. Research Centre for Prevention and Health, Capital Region of Denmark, Copenhagen, Denmark  
227 122. Department of Clinical Experimental Research, Glostrup University Hospital, Glostrup, Denmark  
228 123. Department of Clinical Medicine, Faculty of Health and Medical Sciences, University of Copenhagen,  
229 Copenhagen, Denmark  
230 124. Dasman Diabetes Institute, Dasman, Kuwait  
231 125. Centre for Vascular Prevention, Danube-University Krems, Krems, Austria  
232 126. King Abdulaziz University, Jeddah, Saudi Arabia  
233 127. School of Molecular, Genetic and Population Health Sciences, University of Edinburgh, Medical  
234 School, Teviot Place, Edinburgh, UK  
235 128. HUNT Research Centre, Department of Public Health and General Practice, Norwegian University of  
236 Science and Technology, Levanger, Norway  
237 129. St. Olav Hospital, Trondheim University Hospital, Trondheim, Norway

238 130. Department of Medicine, Levanger Hospital, Nord- Trøndelag Health Trust, Levanger, Norway  
239 131. Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor,  
240 Michigan, USA  
241 132. Department of Human Genetics, University of Michigan, Ann Arbor, Michigan, USA  
242 133. Folkhälsan Research Centre, Helsinki, Finland; Department of Endocrinology, Helsinki University  
243 Central Hospital, Helsinki, Finland  
244 134. Institute for Molecular Medicine Finland University of Helsinki, Helsinki, Finland  
245 135. Department of Clinical Sciences, Diabetes and Endocrinology, Lund University Diabetes Centre,  
246 Malmö, Sweden.  
247 136. Department of Primary Health Care, Vaasa Central Hospital, Vaasa, Finland  
248 137. Diabetes Center, Vaasa Health Care Center, Vaasa, Finland  
249 138. Psychiatric and Neurodevelopmental Genetics Unit, Department of Psychiatry, Massachusetts General  
250 Hospital, Boston, Massachusetts, USA  
251 139. Department of Public Health, University of Helsinki, Finland  
252 140. ICDDR, B; Mohakhali, Dhaka, Bangladesh  
253 141. National Institute of Cardiovascular Diseases, Sher-e-Bangla Nagar, Dhaka, Bangladesh  
254 142. International Centre for Circulatory Health, Imperial College London, UK  
255 143. Molecular and Cellular Therapeutics, Royal College of Surgeons in Ireland, Dublin, Ireland  
256 144. University of Lille, UMR1167, Risk Factors and Molecular Determinants of aging-related diseases,  
257 Lille, France  
258 145. Inserm, Lille, France  
259 146. Centre Hospitalier Universitaire Lille, Public Health, Lille, France  
260 147. Institut Pasteur de Lille, Lille, France  
261 148. Department of Epidemiology and Public Health, EA 3430, University of Strasbourg, Strasbourg,  
262 France  
263 149. Department of General and Interventional Cardiology, University Heart Center Hamburg, Germany  
264 150. University Medical Center Hamburg-Eppendorf, Hamburg, Germany  
265 151. Department of Epidemiology, UMR 1027- INSERM, Toulouse University-CHU Toulouse, Toulouse,  
266 France  
267 152. Director, UKCRC Centre of Excellence for Public Health, Queens University, Belfast, Northern Ireland  
268 153. Institute of Genetic Epidemiology, Helmholtz Zentrum München - German Research Center for  
269 Environmental Health, Neuherberg, Germany  
270 154. Department of Medicine I, University Hospital Grosshadern, Ludwig-Maximilians-Universität,  
271 Munich, Germany  
272 155. DZHK (German Centre for Cardiovascular Research), partner site Munich Heart Alliance, Munich,  
273 Germany  
274 156. Research Center in Epidemiology and Preventive Medicine, Department of Clinical and Experimental  
275 Medicine, University of Insubria, Varese, Italy  
276 157. Princess Al-Jawhara Al-Brahim Centre of Excellence in Research of Hereditary Disorders (PACER-  
277 HD), King Abdulaziz University, Jeddah, Saudi Arabia  
278 158. Cardiovascular Research Center, Massachusetts General Hospital, Boston, Massachusetts, USA  
279 159. Department of Medicine, Harvard Medical School, Boston, Massachusetts, USA  
280 160. A Collaboration between the University Medical Schools and NHS, Aberdeen, Dundee, Edinburgh and  
281 Glasgow, UK  
282 161. Department of Genetics, University of North Carolina, Chapel Hill, North Carolina, USA  
283 162. Institute of Cardiovascular Sciences, University of Manchester, Manchester, UK  
284 163. Faculty of Population Health Sciences, Institute of Cardiovascular Science, University College London,  
285 London, UK

286 164. The Big Data Institute at the Li Ka Shing Centre for Health Information and Discovery, University of  
287 Oxford, Oxford, UK  
288 165. Wellcome Trust Sanger Institute, Hinxton, UK  
289 166. The National Institute for Health Research Blood and Transplant Research  
  
290 167. Contributed equally to this work  
291 168. Jointly supervised the work  
292  
293 Corresponding authors: Joanna M M Howson, [jmmh2@medschl.cam.ac.uk](mailto:jmmh2@medschl.cam.ac.uk) and Patricia B Munroe,  
294 [p.b.munroe@qmul.ac.uk](mailto:p.b.munroe@qmul.ac.uk)  
295  
296  
297



## Abstract

High blood pressure is a major risk factor for cardiovascular disease and premature death. However, there is limited knowledge on specific causal genes and pathways. To better understand the genetics of blood pressure, we genotyped 242,296 rare, low-frequency and common genetic variants in up to ~192,000 individuals, and used ~155,063 samples for independent replication. We identified 31 novel blood pressure or hypertension associated genetic regions in the general population, including three rare missense variants in *RBM47*, *COL21A1* and *RRAS* with larger effects ( $>1.5\text{mmHg/allele}$ ) than common variants. Multiple rare, nonsense and missense variant associations were found in *A2ML1* and a low-frequency nonsense variant in *ENPEP* was identified. Our data extend the spectrum of allelic variation underlying blood pressure traits and hypertension, provide new insights into the pathophysiology of hypertension and indicate new targets for clinical intervention.

## 310 **Introduction**

311 High blood pressure (BP) or hypertension is a highly prevalent chronic disorder. It is estimated to be  
312 responsible for a larger proportion of global disease burden and premature mortality than any other disease  
313 risk factor<sup>1</sup>. Elevated systolic and/or diastolic BP increases the risk of several cardiovascular disorders  
314 including stroke, coronary heart disease (CHD), heart failure, peripheral arterial disease and abdominal  
315 aortic aneurysms<sup>2</sup>. BP is a complex, heritable, polygenic phenotype for which genome-wide association  
316 studies (GWAS) have identified over 67 genetic regions associated with BP and/or hypertension to date<sup>3-11</sup>.  
317 These variants are common (minor allele frequency,  $MAF \geq 0.05$ ), mostly map to intronic or intergenic  
318 regions, with the causal alleles and genes not readily identified due to linkage disequilibrium (LD)<sup>4,5</sup>, and  
319 explain only ~2% of trait variance<sup>12</sup>. Low-frequency ( $0.01 < MAF < 0.05$ ) and rare ( $MAF \leq 0.01$ ) single  
320 nucleotide variants (SNVs), predominantly unexplored by GWAS may have larger phenotypic effects than  
321 common SNVs<sup>13</sup>, and may help to explain the missing heritability, and identify causative genes as  
322 demonstrated previously<sup>14</sup>.

323 To identify novel coding variants and loci influencing BP traits and hypertension we performed the largest  
324 meta-analysis to date that included a total of ~350,000 individuals, directly genotyped with the Exome chip.  
325 The Exome chip contains ~240,000 mostly rare and low-frequency variants (Methods). A single-variant  
326 discovery analysis was performed, and candidate SNVs were taken forward for validation using independent  
327 replication samples. Gene-based tests were used to identify BP associated genes harboring multiple rare  
328 variant associations. We next assessed whether the newly identified BP associated SNVs were associated  
329 with expression levels of nearby genes, and tested these variants in aggregate for a causal association of BP  
330 with other cardiovascular traits and risk factors. Our findings highlight the contribution of rare variants in  
331 the aetiology of blood pressure in the general population, and provide new insights into the pathophysiology  
332 of hypertension.

333

## 334 **Results**

### 335 **Discovery of single variant BP associations**

336 We genotyped 192,763 individuals from 51 studies, and assessed association of 242,296 SNVs with  
337 diastolic BP (DBP), systolic BP (SBP), pulse pressure (PP) and hypertension (HTN; Supplementary Tables  
338 1, 2 and 3; Methods). An overview of the SNV discovery study design is given in Figure 1. A fixed effects  
339 meta-analysis for each trait was performed using study-level association summary statistics from i) samples  
340 of European (EUR) ancestry (up to 165,276 individuals), and ii) a trans-ethnic meta-analysis of the EUR and  
341 additional South Asian (SAS) ancestry samples (EUR\_SAS; up to 192,763 individuals). Two analyses of  
342 DBP, SBP and PP were performed, one in which the trait was inverse normal transformed and a second in  
343 which the raw phenotype was analysed. Both sets of results were consistent (Methods), therefore to  
344 minimise sensitivity to deviations from normality in the analysis of rare variants, the results from the  
345 analyses of the transformed traits were used for discovery. Strong correlations between the BP traits were  
346 observed across studies (Methods), hence no adjustment of significance thresholds for independent trait  
347 testing was applied.

348 The discovery meta-analyses identified 50 genomic regions with genome-wide significant (GWS) evidence  
349 of association with at least one of the four BP traits tested ( $P < 5 \times 10^{-8}$ ; Supplementary Table 4). There were  
350 45 regions associated in the EUR\_SAS samples, of which 13 were novel (Figure 2). An additional five  
351 regions were GWS in the EUR only meta-analyses of which two were novel (Supplementary Figure 1). In  
352 total, 16 genomic regions were identified that were GWS for at least one BP trait that have not been  
353 previously reported.

### 354 **Replication of single variant BP associations**

355 Next we sought support for our findings, in an independent replication dataset comprising of 18 studies, 16  
356 of which were from the Cohorts for Heart and Aging Research in Genomic Epidemiology+ (CHARGE+)  
357 exome chip blood pressure consortium (Figure 1; Liu *et al.* Nature Genetics, *submitted*). Variants were  
358 selected for replication first using the larger (transformed) EUR\_SAS data, with additional variants from the  
359 (transformed) EUR data also selected. SNVs were selected if they mapped outside of known BP genomic  
360 regions and had  $MAF \geq 0.05$  and  $P < 1 \times 10^{-5}$  or  $MAF < 0.05$  and  $P < 1 \times 10^{-4}$  with at least one BP trait, *i.e.*  
361 choosing a lower significance threshold for the selection of rare variants (full details of the selection criteria  
362 are provided in the Methods). In total 81 candidate SNVs were selected for replication (Supplementary

Table 5). Eighty variants were selected from EUR\_SAS (transformed) results and one SNV at the *ZNF101* locus from the EUR (transformed) analyses. The results for EUR\_SAS and EUR were consistent (association statistics were correlated,  $\rho=0.9$  across ancestries for each of the traits). Of the 81 variants, 30 SNVs were selected for association with DBP as the primary trait, 26 for SBP, 19 for PP and 6 for HTN, with the primary trait defined as the BP trait with the smallest association *P*-value in the EUR-SAS discovery analyses.

Meta-analyses were performed on results from analyses of untransformed DBP, SBP, PP and HTN (as only results of untransformed traits were available from CHARGE+) in (i) up to 125,713 individuals of EUR descent, and (ii) up to 155,063 individuals of multiple ethnicities (4,632 of Hispanic descent, 22,077 of African American descent, 2,641 SAS samples with the remainder EUR; Figure 1). Given that a large proportion of the ancestries in the trans-ethnic meta-analyses were not included in our discovery samples, we used the EUR meta-analyses as the main data set for replication, but we also report any additional associations identified within the larger trans-ethnic dataset.

Novel BP-SNV associations were identified based on two criteria (Figure 1; Methods). Firstly, replication of the primary BP trait-SNV association was sought at a Bonferroni adjusted *P*-value threshold in the replication data ( $P \leq 6.17 \times 10^{-4}$ , assuming  $\alpha=0.05$  for 81 SNVs tested and same direction of effect; Methods) without the need for GWS. Secondly, meta-analyses of discovery and replication results across all four (untransformed) BP traits were performed to assess the overall level of support across all samples for the 81 candidate SNVs; those BP-SNV associations that were GWS (with statistical support in the replication studies;  $P < 0.05$  and the same direction of effect) were also declared as novel.

Seventeen SNV-BP associations formally replicated with concordant direction of effect at a Bonferroni adjusted significance level for the primary trait. Fourteen were in the EUR meta-analyses, and amongst these was a rare non-synonymous (ns) SNV mapping to *COL21A1* (Table 1, Supplementary Table 6). Three associations were in the trans-ethnic meta-analyses, these included two rare nsSNVs in *RBM47* and *RRAS* (Table 1, Supplementary Table 7; Methods).

389

390 In addition to the 17 SNV-BP trait associations that formally replicated, we identified 13 further SNV-  
 391 associations that were GWS in the combined (discovery and replication) meta-analyses. Ten of these were  
 392 GWS in the combined EUR analyses, (Table 2; Supplementary Tables 6 and 8a), and three were GWS in the  
 393 combined trans-ethnic meta-analyses (Table 2; Supplementary Tables 7 and 8b).

394

395 This gives a total of 30 novel SNV-BP associations (15 SNV-DBP, 9 SNV-SBP and 6 SNV-PP; Tables 1  
 396 and 2; Supplementary Figures 2 and 3). Five of the SNVs were GWS with more than one BP trait (Figure 3:  
 397 Tables 1 and 2; Supplementary Table 8). Four loci (*CERS5*, *TBX2*, *RGL3* and *OBFC1*) had GWS  
 398 associations with HTN in addition to GWS associations with DBP and SBP. The *PRKAG1* locus had GWS  
 399 associations with both SBP and PP.

400

401 Conditional analyses were performed to identify secondary signals of association within the novel BP loci.  
 402 The RAREMETALWORKER (RMW) package (Methods)<sup>15</sup> allows conditional analyses to be performed  
 403 using summary level data. Hence, analyses of the transformed primary traits and HTN were re-run in RMW  
 404 across the discovery studies (Figure 4). The results of the RMW single variant tests were consistent with the  
 405 initial discovery analyses (Supplementary Information). Given the RMW analyses were based on our  
 406 discovery samples, the larger EUR-SAS data was used as the main analysis to increase power, but we also  
 407 report any additional associations with evidence in EUR.

408

409 We identified secondary independent signals of association in four loci, *PREX1*, *PRKAG1* and *RRP1B*  
 410 within the EUR\_SAS analyses and *COL21A1* in the EUR analyses ( $P_{\text{conditional}} < 1 \times 10^{-4}$ , Bonferroni adjusted  
 411 for ~500 variants within each region; Methods; Supplementary Tables 9 and 10). Three independent  
 412 association signals were identified in the *MYH6* locus in the EUR\_SAS analyses (Supplementary Table 11).

413

#### 414 **Gene-based BP associations**

415 To improve statistical power to detect associations in genes harbouring rare variants, analytical methods that  
 416 combine effects of variants across a gene into a single test have been devised and are implemented in the

RMW package<sup>15</sup>. We applied the gene-based sequence kernel association test (SKAT)<sup>16</sup> and Burden tests<sup>17</sup> to the RMW dataset (MAF<0.05 or MAF<0.01; Figure 4; Methods). One previously unidentified BP gene (*A2ML1*) was associated with HTN ( $P=7.73 \times 10^{-7}$ ) in the EUR\_SAS studies and also in EUR studies (Supplementary Table 12; Bonferroni-corrected threshold of significance  $P < 2.8 \times 10^{-6}$ , after adjusting for 17,996 genes tested, Methods). The gene showed residual association with the primary BP trait after conditioning on the most associated SNV in the gene ( $P_{\text{conditional}} = 5.00 \times 10^{-4}$ ; Supplementary Table 12), suggesting that the association is due to multiple rare variants in the gene. One nonsense variant (rs199651558, p.Arg893\*, MAF= $3.5 \times 10^{-4}$ ) was observed, and there were multiple missense variants (Figure 5). *A2ML1* encodes alpha-2-macroglobulin-like 1 protein, and is a member of the alpha macroglobulin superfamily, which comprises protease inhibitors targeting a wide range of substrates. Mutations in this gene are associated with a disorder clinically related to Noonan syndrome, a developmental disorder which involves cardiac abnormalities<sup>18</sup>. We sought replication in the CHARGE+ studies for this gene, however there was no evidence of association with HTN ( $P=0.45$ ). Given the very low frequencies of the variants involved, however, studies in which the variants are polymorphic will be required to replicate the association with HTN. The DBH gene was found to be associated with DBP using the SKAT test ( $P=2.88 \times 10^{-6}$ ). However, this was not due to multiple rare variants as the association was driven by rs77273740 (Supplementary Table 5) and the SNV was not validated in the replication samples.

### Rare and common variant associations in established BP loci

Of the 67 established BP loci, 35 loci were on the Exome chip (N=43 SNVs or close proxies  $r^2 > 0.7$ ). All 43 SNVs had at least nominal evidence of association with BP in our discovery samples ( $P < 0.01$ ; Supplementary Table 13). We also assessed if any of the established BP loci contained coding variants that are associated with BP traits and in LD ( $r^2 > 0.2$ ) with the known BP variants on the Exome chip (Supplementary Table 13), using the 1000G phase 3 release for LD annotation. Focusing on SNVs that were GWS for any BP trait from our transformed discovery data for either ancestry, there were 25 coding variants, of which 6 were predicted to be damaging at loci labelled *CDC25A*, *SLC39A8*, *HFE*, *ULK4*, *ST7L-CAPZA1-MOV10* and *CYP11A1-ULK3*. Three of these are published variants at loci labelled *SLC39A8*, *HFE* and *ST7-CAPZA1-MOV10*. At *CYP11A1-ULK3*, the coding variant was in moderate LD with the reported

variant, but was less significantly associated with DBP in our EUR\_SAS dataset ( $P=2.24 \times 10^{-8}$  compared to  $P=1.68 \times 10^{-15}$  for the published variant). At the *ULK4* locus the predicted damaging coding variant had similar association as the published coding variant (predicted to be benign), and prior work has already indicated several associated nsSNVs in strong LD in *ULK4*<sup>19</sup>. The nsSNV within the *CDC25A* locus (rs11718350 in *SPINK8*) had similar association with DBP as the intergenic published SNV in our EUR\_SAS dataset ( $P=2.00 \times 10^{-8}$  compared to  $P=2.27 \times 10^{-8}$  for the published variant). Overall at least 5 of the known loci are consistent with having a coding causal variant.

Gene-based SKAT tests of all genes that map within 1 Mb of a previously reported SNV association (Supplementary Table 14), indicated no genes with multiple rare or low-frequency variant associations. Single variant conditional analyses showed that rs33966350, a rare nonsense variant in *ENPEP* (MAF=0.01) was associated with SBP ( $P_{\text{conditional}}=1.61 \times 10^{-5}$ ) in the EUR\_SAS samples (Supplementary Tables 14 and 15; Methods) independently of the known SNV (rs6825911). *ENPEP* encodes aminopeptidase A (APA) an enzyme of the renin-angiotensin-aldosterone system (RAAS) that converts angiotensin II (AngII) to AngIII. There were no other established loci with convincing low-frequency or rare SNV associations in the EUR\_SAS samples. However, *HOXC4*, had evidence of a second independent signal with a rare missense SNV in EUR samples (rs78731604; MAF=0.005,  $P_{\text{conditional}}=5.76 \times 10^{-5}$ ; Supplementary Table 15). The secondary signal in the *HOXC4* region, mapped to *CALCOCO1*, ~300kb from the known SNV. The gene association (MAF $\leq$ 0.01,  $P=2.37 \times 10^{-5}$ ) was below the required significance threshold and attributable to rs78731604, which is not predicted to have detrimental effects on protein structure. Therefore, replication of this association is required. Three loci (*ST7L-CAPZA1-MOV10*, *FIGN-GRB14*, and *TBX5-TBX3*) had evidence of a second independent signal in the region in EUR\_SAS samples with a common variant ( $P_{\text{conditional}} < 1 \times 10^{-4}$ ; Supplementary Table 15) that has not been previously reported.

Having identified 30 novel loci associated with BP traits, as well as additional new independent SNVs at four novel loci and five known loci, we calculated the percent of the trait variance explained (Methods). This was 2.08%/2.11%/1.15% for SBP/DBP/PP for the 43 previously reported BP-SNVs covered in our dataset, increasing to 3.38%/3.41%/2.08% respectively with the inclusion of the 30 lead SNVs from novel loci, plus new independent SNV-BP associations identified from novel and known loci.

473 **Effect of BP SNVs on cardiovascular traits & risk factors**

474 Amongst our novel BP-SNV associations, some have previously been reported to be associated with other  
 475 cardiovascular traits and risk factors (Supplementary Table 16); these include coronary heart disease (CHD:  
 476 *PHACTR1*, *ABO*)<sup>20,21</sup>, QT interval (*RNF207*)<sup>22</sup>, heart rate (*MYH6*)<sup>23</sup>, and cholesterol levels (2q36.3, *ABO*,  
 477 *ZNF101*)<sup>24</sup>.

478 To test the impact of BP variants on cardiovascular endpoints and risk factors we created three weighted  
 479 genetic risk scores (GRS) according to SBP/DBP/PP based on the newly identified and previously published  
 480 BP variants (up to N=125; Methods). The GRS models were used to test the causal effect of BP on the  
 481 following traits: ischemic stroke (including the subtypes, cardiometabolic, large and small vessel <sup>25</sup>), CHD,  
 482 heart failure,<sup>26</sup> left ventricular mass<sup>27</sup>, left ventricular wall thickness<sup>27</sup>, high-density lipoprotein cholesterol  
 483 (HDL-c), low-density lipoprotein (LDL-c), triglycerides, total cholesterol, body mass index (BMI), waist-  
 484 hip ratio adjusted BMI, height and estimated glomerular filtration rate (eGFR) (Methods). As expected, BP  
 485 was positively associated with increased CHD risk (OR [95% CI]=1.39[1.22-1.59] per 10mmHg increase in  
 486 SBP,  $P=6.07 \times 10^{-7}$ ; 1.62[1.28-2.05] per 10mmHg increase in DBP,  $P=5.99 \times 10^{-5}$ ; 1.70[1.34-2.16] per  
 487 10mmHg increase in PP,  $P=1.20 \times 10^{-5}$ ; Table 3), and increased risk of ischemic stroke (OR [95%  
 488 CI]=1.93[1.47-2.55] per 10mmHg increase in DBP,  $P=2.81 \times 10^{-6}$ ; 1.57[1.35-1.84] per 10mmHg increase in  
 489 SBP,  $P=1.16 \times 10^{-8}$ ; 2.12[1.58-2.84] per 10mmHg increase in PP,  $P=5.35 \times 10^{-7}$ ). The positive association with  
 490 ischemic stroke was primarily due to large vessel stroke (Table 3). DBP and SBP were also positively  
 491 associated with left ventricular mass (9.57 [3.98-15.17] gram increase per 10mmHg increase in DBP,  
 492  $P=8.02 \times 10^{-4}$  and 5.13 [1.77-8.48] gram increase per 10mmHg increase in SBP,  $P=0.0027$ ) and left  
 493 ventricular wall thickness (0.10 [0.06-0.13] cm increase per 10mmHg increase in DBP,  $P=1.88 \times 10^{-8}$  and  
 494 0.05 [0.03-0.07] cm increase per 10mmHg increase in SBP,  $P=5.52 \times 10^{-6}$ , Table 3). There was no convincing  
 495 evidence to support the BP associated variants having an effect on lipid levels ( $P>0.1$ ), BMI ( $P>0.005$ ),  
 496 waist hip ratio adjusted BMI ( $P>0.1$ ), height ( $P>0.06$ ), eGFR ( $P>0.02$ ) or heart failure ( $P>0.04$ ). The causal  
 497 associations with CHD, stroke, and left ventricular measures augment the results from a previous association  
 498 analysis using 29 BP variants<sup>28</sup>. Our data strongly support the previous observations of no causal



relationship between BP and eGFR. Lack of evidence of a BP effect with heart failure may only be due to lack of power, as the association was in the expected direction.

### Possible functional variants at BP loci and candidate genes

Twenty-six of our newly discovered BP associated SNVs had  $MAF \geq 0.05$  and therefore due to extensive LD with other SNVs not genotyped on the Exome array, identifying the causal genes requires additional information. If a SNV is associated with increased or decreased expression of a particular gene, *i.e.* it is an expression quantitative trait locus (eQTL) this suggests the gene on which the SNV acts could be in the causal pathway. To help identify potential candidate causal genes in the novel BP loci (Supplementary Table 9), information from publicly available eQTL databases was investigated (MuTHER for LCL, adipose and skin and GTEx for nine tissues including the heart and tibial artery; Methods).

The DBP increasing allele of the nsSNV, rs7302981-A, was associated with increased expression of *CERS5* in: LCLs ( $P_{MuTHER}=3.13 \times 10^{-72}$ ) skin ( $P_{MuTHER}=2.40 \times 10^{-58}$ ) adipose ( $P_{MuTHER}=2.87 \times 10^{-54}$ ) and nerve ( $P_{GTEx}=4.5 \times 10^{-12}$ ) (Supplementary Figure 4). Additional testing (Methods) provided no evidence against colocalisation of the eQTL and DBP association signals, implicating *CERS5* as a candidate causal gene for this DBP locus. *CERS5* (LAG1 homolog, ceramide synthase 5) is involved in the synthesis of ceramide, a lipid molecule involved in several cellular signaling pathways. *Cers5* knockdown has been shown to reduce cardiomyocyte hypertrophy in mouse models<sup>29</sup>. However, it is unclear whether the blood pressure raising effects at this locus are the cause or result of any potential effects on cardiac hypertrophy. Future studies investigating this locus in relation to parameters of cardiac hypertrophy and function (*e.g.* ventricular wall thickness) should help address this question.

The DBP raising allele of the nsSNV (rs867186-A) was associated with increased expression of *PROCR* in adipose tissue ( $P_{MuTHER}=3.24 \times 10^{-15}$ ) and skin ( $P_{MuTHER}=1.01 \times 10^{-11}$ ) (Supplementary Figure 4). There was no evidence against colocalisation of the eQTL and DBP association thus supporting *PROCR* as a candidate causal gene. *PROCR* encodes the Endothelial Protein C receptor, a serine protease involved in the blood

coagulation pathway, and rs867186 has previously been associated with coagulation and haematological factors.<sup>30,31</sup> The PP decreasing allele of, rs10407022-T, which is predicted to have detrimental effects on protein structure (Methods) was associated with increased expression of *AMH* in muscle ( $P_{\text{GTEX}}=9.95 \times 10^{-15}$ ), thyroid ( $P_{\text{GTEX}}=8.54 \times 10^{-7}$ ), nerve ( $P_{\text{GTEX}}=7.15 \times 10^{-8}$ ), tibial artery ( $P_{\text{GTEX}}=6.46 \times 10^{-9}$ ), adipose ( $P_{\text{GTEX}}=4.69 \times 10^{-7}$ ), and skin ( $P_{\text{GTEX}}=5.88 \times 10^{-8}$ ) (Supplementary Figure 4). There was no evidence against colocalisation of the eQTL and PP association, which supports *AMH* as a candidate causal gene for PP. Low *AMH* levels have been previously associated with hypertensive status in women with the protein acting as a marker of ovarian reserve<sup>32</sup>. The intergenic SBP raising allele of rs4728142-A was associated with reduced expression of *IRF5* in skin ( $P_{\text{MUTHER}}=5.24 \times 10^{-31}$ ) and LCLs ( $P_{\text{MUTHER}}=1.39 \times 10^{-34}$ ), whole blood ( $P_{\text{GTEX}}=3.12 \times 10^{-7}$ ) and tibial artery ( $P_{\text{GTEX}}=1.71 \times 10^{-7}$ ).

535

Three novel rare nsSNVs were identified that map to *RBM47*, *RRAS* (both associated with SBP) and *COL21A1* (associated with PP). They had larger effect sizes than common variant associations ( $>1.5$  mmHg per allele; Supplementary Figure 5) and were predicted to have detrimental effects on protein structure (Supplementary Table 16; Methods). In *RBM47*, rs35529250 (p.Gly538Arg) is located in a highly conserved region of the gene and was most strongly associated with SBP (MAF=0.008; +1.59 mmHg per T allele;  $P=5.90 \times 10^{-9}$ ). *RBM47* encodes the RNA binding motif protein 47 and is responsible for post-transcriptional regulation of RNA, through its direct and selective binding with the molecule.<sup>33</sup> In *RRAS*, rs61760904 (p.Asp133Asn) was most strongly associated with SBP (MAF=0.007; +1.51 mmHg per T allele;  $P=8.45 \times 10^{-8}$ ). *RRAS* encodes a small GTPase belonging to the Ras subfamily of proteins H-RAS, N-RAS, and K-RAS and has been implicated in actin cytoskeleton remodelling, and controlling cell proliferation, migration and cycle processes<sup>34</sup>. The nsSNV in *COL21A1* (rs200999181, p.Gly665Val) was most strongly associated with PP (MAF=0.001; +3.14 mmHg per A allele;  $P=1.93 \times 10^{-9}$ ). *COL21A1* encodes the collagen alpha-1 chain precursor of type XXI collagen, a member of the FACIT (fibril-associated collagens with an interrupted triple helix) family of proteins<sup>35</sup>. The gene is detected in many tissues, including the heart and aorta. Based on our results, these three genes represent good candidates for functional follow-up. However, due to the incomplete coverage of all SNVs across the region on the Exome chip, it is possible that other non-genotyped SNVs may better explain some of these associations. We therefore checked for variants in LD

553 ( $r^2 > 0.3$ ) with these three rare nsSNVs in the UK10K + 1000G dataset<sup>36</sup> to ascertain if there are other  
554 candidate SNVs at these loci (Supplementary Table 17). There were no SNVs within 1Mb of the *RBM47*  
555 locus in LD with the BP associated SNV. At the *COL21A1* locus there were only SNVs in moderate LD, and  
556 these were annotated as intronic, intergenic or in the 5'UTR. At the *RRAS* locus, there were two SNVs in  
557 strong LD with the BP associated SNV, which both mapped to introns of *SCAF1* and are not predicted to be  
558 damaging. All SNVs in LD at both loci were rare as expected (Supplementary Table 17) supporting a role  
559 for rare variants. Hence, the rare BP associated nsSNVs at *RBM47*, *COL21A1* and *RRAS* remain the best  
560 causal candidates.

## 561

### 562 **Pathway and network analyses**

563 To identify connected gene sets and pathways implicated by the BP associated genes we used Meta-Analysis  
564 Gene-set Enrichment of variant Associations (MAGENTA)<sup>37</sup> and GeneGo MetaCore (Thomson Reuters,  
565 UK). MAGENTA tests for over-representation of BP associated genes in pre-annotated pathways (gene  
566 sets) (Methods and Supplementary Table 18a). GeneGo Metacore identifies potential gene networks. The  
567 MAGENTA analysis was used for hypothesis generation and results were compared with the GeneGo  
568 Metacore outputs to cross-validate findings.

569 Using MAGENTA there was an enrichment ( $P < 0.01$  and  $FDR < 5\%$  in either the EUR\_SAS or the EUR  
570 participants) of six gene sets with DBP, three gene sets with HTN and two gene sets for SBP  
571 (Supplementary Table 18b). The RNA polymerase I promoter clearance (chromatin modification) pathway  
572 showed the most evidence of enrichment with genes associated with DBP ( $P_{\text{Reactome}} = 8.4 \times 10^{-5}$ ,  $FDR = 2.48\%$ ).  
573 NOTCH signalling was the most associated pathway with SBP ( $P_{\text{Reactome}} = 3.00 \times 10^{-4}$ ,  $FDR = 5\%$ ) driven by  
574 associations at the *FURIN* gene. The inorganic cation anion solute carrier (SLC) transporter pathway had  
575 the most evidence of enrichment by HTN associated genes ( $P_{\text{Reactome}} = 8.00 \times 10^{-6}$ ,  $FDR = 2.13\%$ ).

576 Using GeneGo MetaCore, five network processes were enriched ( $FDR < 5\%$ ; Methods; Supplementary  
577 Tables 19 and 20). These included several networks with genes known to influence vascular tone and BP:  
578 inflammation signalling,  $P = 1.14 \times 10^{-4}$  and blood vessel development  $P = 2.34 \times 10^{-4}$ . The transcription and  
579 chromatin modification network ( $P = 2.85 \times 10^{-4}$ ) was also enriched, a pathway that was also highlighted in the

MAGENTA analysis, with overlap of the same histone genes (*HIST1H4C*, *HIST1H2AC*, *HIST1H2BC*, *HIST1H1T*) and has also been recently reported in an integrative network analysis of published BP loci and whole blood expression profiling<sup>38</sup>. Two cardiac development pathways were enriched: the oxidative stress-driven (ROS/NADPH) ( $P=4.12 \times 10^{-4}$ ) and the Wnt/ $\beta$ -catenin/integrin-driven ( $P=0.0010$ ). Both these cardiac development pathways include the *MYH6*, *MYH7*, and *TBX2* genes, revealing a potential overlap with cardiomyopathies and hypertension, and suggesting some similarity in the underlying biological mechanisms.

587

## 588 Discussion

By conducting the largest ever genetic study of BP, we identified further novel common variants with small effects on BP traits, similar to what has been observed for obesity and height<sup>39,40</sup>. More importantly, our study identified some of the first rare coding variants of strong effect ( $>1.5\text{mmHg}$ ) that are robustly associated with BP traits in the general population, complementing and extending the previous discovery and characterisation of variants underlying rare Mendelian disorders of blood pressure regulation<sup>41</sup>. Using SNV associations in 17 genes reported to be associated with monogenic disorders of blood pressure (Methods) we found no convincing evidence of enrichment ( $P_{\text{enrichment}}=0.044$ ). This suggests that BP control in the general population may occur through different pathways to monogenic disorders of BP re-enforcing the importance of our study findings. The identification of 30 novel BP loci plus further new independent secondary signals within four novel and five known loci (Methods) has augmented the trait variance explained by 1.3%, 1.2% and 0.93% for SBP, DBP and PP respectively within our data-set. This suggests that with substantially larger sample sizes, for example through UK BioBank<sup>42</sup>, we expect to identify 1000s more loci associated with BP traits, and replicate more of our discovery SNV associations that are not yet validated in the current report.

The discovery of rare missense variants has implicated several interesting candidate genes, which are often difficult to identify from common variant GWAS, and should therefore lead to more rapidly actionable biology. *A2ML1*, *COL21A1*, *RRAS* and *RBM47* all warrant further follow-up studies to define the role of

606 these genes in regulation of BP traits, as well as functional studies to understand their mechanisms of action.  
607 *COL21A1* and *RRAS* warrant particular interest since both are involved in blood vessel remodelling, a  
608 pathway of known aetiological relevance to hypertension.

609 We observed a rare nonsense SBP associated variant in *ENPEP* (rs33966350; p.Trp317\* ): this overlaps a  
610 highly conserved region of both the gene and protein and is predicted to result in either a truncated protein  
611 with reduced catalytic function or is subject to nonsense mediated RNA decay. ENPEP converts angiotensin  
612 II (AngII) to Ang-III. AngII activates the angiotensin 1 (AT1) receptor resulting in vasoconstriction, while  
613 AngIII activates the angiotensin 2 (AT2) receptor that promotes vasodilation and protects against  
614 hypertension.<sup>43</sup> The predicted truncated protein may lead to predominant AngII signaling in the body, and  
615 increases in BP. This new observation could potentially inform therapeutic strategies. Of note, angiotensin-  
616 converting-enzyme (ACE) inhibitors are commonly used in the treatment of hypertension. However, patients  
617 who suffer from adverse reactions to ACE inhibitors, such as dry cough and skin rash, would benefit from  
618 alternative drugs that target RAAS. Murine studies have shown that in the brain, AngIII is the preferred AT1  
619 agonist that promotes vasoconstriction and increases blood pressure, as opposed to AngII in the peripheral  
620 system. These results have motivated the development of brain specific APA inhibitors to treat  
621 hypertension<sup>44</sup>. Our results confirm APAs, such as ENPEP, as a valid target to modify blood pressure, but  
622 suggest that long-term systemic reduction in APA activity may lead to an increase in blood pressure. Future  
623 studies are needed to examine the effects of the p.Trp317\* variant on the RAAS system, specifically in the  
624 brain and peripheral vasculature, in order to test the benefits of the proposed therapeutic strategy in humans.

625 In addition to highlighting new genes in pathways of established relevance to BP and hypertension, and  
626 identifying new pathways, we have also identified multiple signals at new loci. For example, there are three  
627 distinct signals at the locus containing the *MYH6/MYH7* genes, and we note that *TBX2* maps to one of the  
628 novel regions. These genes are related to cardiac development and/or cardiomyopathies, and provide an  
629 insight into the shared inheritance of multiple complex traits. Unravelling the causal networks within these  
630 polygenic pathways may provide opportunities for novel therapies to treat or prevent both hypertension and  
631 cardiomyopathies.

## 633 **URLs**

634 Exome chip design information: [http://genome.sph.umich.edu/wiki/Exome\\_Chip\\_Design](http://genome.sph.umich.edu/wiki/Exome_Chip_Design)

635 [RareMetalWorker](http://genome.sph.umich.edu/wiki/RAREMETALWORKER) information: <http://genome.sph.umich.edu/wiki/RAREMETALWORKER>

636 [Summary SNV association results: http://www.phenoscanter.medschl.cam.ac.uk](http://www.phenoscanter.medschl.cam.ac.uk)

637 [Databases used for variant annotation: http://www.ncbi.nlm.nih.gov/SNP/](http://www.ncbi.nlm.nih.gov/SNP/)

638 <http://www.ensembl.org/info/docs/tools/index.html> and <http://evs.gs.washington.edu/EVS/>

639 UCSC reference file used for annotation of variants with gene and exon information:

640 <http://hgdownload.soe.ucsc.edu/goldenPath/hg19/database/refFlat.txt.gz>

641 Databases used for pathway analysis: MAGENTA (<https://www.broadinstitute.org/mpg/magenta/>) and

642 THOMSON REUTERS MetaCore™ Single Experiment Analysis workflow tool

643 ([http://thomsonreuters.com/en/products-services/pharma-life-sciences/pharmaceutical-](http://thomsonreuters.com/en/products-services/pharma-life-sciences/pharmaceutical-research/metacore.html)

644 [research/metacore.html](http://thomsonreuters.com/en/products-services/pharma-life-sciences/pharmaceutical-research/metacore.html)).

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## Author contributions

Supervision and management of the project: JMMH and PBM. The following authors contributed to the drafting of the manuscript: JMMH, PBM, PSu, HW, ASB, FD, JPC, DRB, KW, MT, FWA, LVW, NJS, JD AKM, HY, CMM, NG, XS, TaT, DFF, MHs, OG, TF, VT. All authors critically reviewed and approved the final version of the manuscript. **Statistical analysis review:** JMMH, PSu, FD, HW, JPC, RY, NM, PBM, LVW, HY, TF, EMi, ADM, AM, AM, EE, ASB, FWA, MJC, CF, TF, SEH, ASH, JEH, JL, GM, JM, NM, APM, APo, NJS, RAS, LS, KE, MT, VT, TVV, NV, KW, AMY, WZg, NG, CML, AKM, XS, TT. **Central Data QC:** JMMH, ASB, PSu, RY, FD, HW, JPC, TF, LVW, PBM, EMi, NM, CML, NG, XS, AKM. **Central Data analysis:** JMMH, PSu, FD, HW, JPC, NG, CML, AKM, XS. **Pathway analysis and**

992 **literature review:** JMMH, DRB, PBM, MT, KW, VT, OG, AT, FWA. **GWAS lookups, eQTL analysis,**  
993 **GRS, variant annotation and enrichment analyses:** JMMH, ASB, DRB, JRS, DFF, FD, MHr, PBM,  
994 FWA, TT, CML, AKM, SBu. **Study Investigators in alphabetical order by consortium (CHD Exome+,**  
995 **ExomeBP and GoT2D):** DSA, PA, EA, DA, ASB, RC, JD, JF, IF, PF, JWJ, FKe, ASM, SFN, BGN, DS,  
996 NSa, JV, FWA, PIWB, MJB, MJC, JCC, JMC, IJD, GD, AFD, PE, TE, PWF, GG, PH, CH, KH, EI, MJ,  
997 FKa, SK, JSK, LLi, MIM, OM, AMe, ADM, APM, PBM, MEN, SP, CP, OPo, DP, SR, OR, IR, VS, NJS,  
998 PSe, TDS, JMS, NJW, CJW, EZ, MB, IB, FSC, LG, TH, EKH, PJ, JKu, ML, TAL, AL, KLM, HO, OPe,  
999 RR, JT, MU. **Study Phenotyping in alphabetical order by consortium (CHD Exome+, ExomeBP and**  
000 **GoT2D):** PA, DA, SBl, MC, JF, JWJ, FKe, KK, SFN, BGN, CJP, AR, MS, NSa, JV, WZo, RAB, MJB,  
001 MJC, JCC, JMC, AFD, ASFD, LAD, TE, AF, GG, GH, PH, AS H, OLH, EI, MJ, FK, JSK, LLi, LLa, GM,  
002 AMc, PM, AMe, RMg, MJN, MEN, OPo, NP, FR, VS, NJS, TDS, AVS, JMS, MT, AV, NV, NJW, TiT,  
003 CC, LLH, MEJ, AK, PK, JL DPS, SM, ERBP, AS, TS, HMS, BT. **Study Data QC and analysis in**  
004 **alphabetical order by consortium (CHD Exome+, ExomeBP and GoT2D):** ASB, AJMC, JMMH, JK,  
005 SFN, BGN, MMN, SP, MP, PSu, ST, GV, SMW, RY, FWA, JPC, FD, AF, TF, CH, AMc, AMj, APM,  
006 PBM, CP, WR, FR, NJS, MT, VT, HW, HY, NG, AKM, XS. **Exome chip data QC in alphabetical order**  
007 **by consortium (CHD Exome+, ExomeBP and GoT2D):** ASB, JMMH, SFN, BGN, PSu, RY, FWA,  
008 PIWB, AIFB, JCC, JPC, PD, LAD, FD, EE, CF, TF, SEH, PH, SSH, KH, JEH, EK, AMj, GM, JM, NM,  
009 EMi, AMo, APM, PBM, CPN, MJN, CP, AP, WR, NRR, RAS, NS, LS, KES, MDT, VT, TVV, TVV, NV,  
010 HW, HY, AMY, EZ, WZg, NG, CML, AKM, XS. **Exome chip Data analysis in alphabetical order by**  
011 **consortium (CHD Exome+, ExomeBP and GoT2D):** JMMH, PSu, RY, FWA, PIWB, AIFB, RAB, MJC,  
012 JCC, JPC, PD, LAD, PE, EE, CF, TF, PWF, SF, CG, SEH, PH, ASH, CH, OLH, JEH, EI, MJ, FKa, JSK,  
013 DCML, LLi, JL, GM, RMr, JM, NM, MIM, PM, OM, CM, EMi, AMo, APM, RMg, PBM, CPN, MJN, TO,  
014 APo, APa, WR, NRR, NJS, RAS, NS, LS, TDS, KES, MDT, ET, VT, TVV, NV, LVW, NJW, HW, HY,  
015 AMY, EZ, HZ, WZg, LLB, APG, NG, MHs, JRH, AUJ, JBJ, CML, AKM, NN, XS, AS, AJS. **GRS**  
016 **lookups:** AEJ, EMa, HFM, HL, HMH, JFF, MTr, RSV, WL.

017

018 **Conflict of interests**

019 N. P. has received financial support from several pharmaceutical companies that manufacture either blood  
020 pressure lowering or lipid lowering agents, or both, and consultancy fees.

021 S. K. has received Research Grant-Merck, Bayer, Aegerion; SAB-Catabasis, Regeneron Genetics Center,  
022 Merck, Celera; Equity-San Therapeutics, Catabasis; Consulting-Novartis, Aegerion, Bristol Myers-Squibb,  
023 Sanofi, AstraZeneca, Alnylam.

024 P. Sever has received research awards from Pfizer Inc.

025 A. Malarstig and M. Uria-Nickelsen are full time employees of Pfizer.

026 D. Reily and M. Hoek are full time employees of Merck and co Inc.

027 M.J. Caulfield is Chief Scientist for Genomics England a UK Government company.

028 The authors declare no competing financial interest.

029

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137 **Figure Legends**

138 **Figure 1 Study design and work flow diagram of single variant discovery analyses.** EUR=European, SAS=South  
139 Asian, HIS=Hispanic, AA=African American, HTN=hypertension, BP=blood pressure, SBP=systolic blood pressure,  
140 DBP= diastolic blood pressure, PP=pulse pressure, N=sample size, MAF=minor allele frequency,  $P$ = $P$ -value  
141 significance threshold, SNV=single-nucleotide variant, GWS=genome-wide significance \*Further details of the selection  
142 criteria are provided in the methods.

143 **Figure 2 Discovery SNV-BP associations.** Results are provided for (a) transformed SBP (b) transformed DBP (c)  
144 transformed PP and (d) HTN in the European and South Asian (EUR\_SAS) discovery samples. The y-axis represents –  
145  $\log_{10}P$  for association. Red triangles represent variants that map to one of the 81 regions selected for replication, blue  
146 triangles represent SNVs that map to previously published BP regions, and grey triangles represent all remaining SNVs.  
147 SNVs are ordered according to chromosome (black lines on the outside of the plot) and physical position. Genes that  
148 SNVs map to are given in the outer blocks.

149  
150 **Figure 3 Overlap of the 30 novel loci associations across SBP, DBP, PP and HTN.** The Venn diagram shows which  
151 of the 30 newly identified BP loci are associated with multiple BP traits. Only SNV-BP trait associations that were  
152 genome-wide significant ( $P < 5 \times 10^{-8}$ ) in the combined discovery and replication meta-analyses are listed for any given  
153 BP trait, within the corresponding ancestry dataset that the given locus was validated for (see Tables 1 and 2). The  
154 association of *RRAS* variant with SBP was replicated in the independent samples, but did not achieve GWS in the  
155 combined discovery and replication meta-analysis and is therefore only included for SBP. HTN=hypertension,  
156 SBP=systolic blood pressure, DBP= diastolic blood pressure, PP=pulse pressure.

157 **Figure 4 Study design for conditional analyses and rare variant gene-based discovery analyses.**  
158 RMW=RareMetalWorker, EUR=European, SAS = South Asian, HTN=hypertension, BP=blood pressure, SBP=systolic  
159 blood pressure, DBP= diastolic blood pressure, PP=pulse pressure. N=sample size, MAF=minor allele frequency,  $P$ = $P$ -  
160 value significance threshold,  $P_{\text{cond}}$ =conditional  $P$ -value significance threshold

161 **Figure 5 Locus plot for *A2ML1* and secondary amino acid structure of the gene product.** (a) Locus plot for *A2ML1*  
162 associated with HTN identified through gene based tests. The variants' positions along the gene (x axis, based on



human genome build 37) and the  $-\log_{10}(P\text{-value of association})$  (y axis) are indicated. The variants are colour coded: nonsense (black), missense, predicted damaging (blue), and missense (orange). The schematic above the x-axis represents the intron / exon (black vertical bars) structure, the untranslated regions are shown as grey vertical bars.

(b) The white box denotes the full-length amino acid sequence for each of the two gene products. Black numbers denote amino acid residue positions of note. Coloured boxes depict putative functional domains (see below). Coloured vertical lines indicate the amino acid substitutions corresponding to the variants depicted in the locus plots above using the same colour coding. Bold, italic indicates the SNV association with smallest *P*-value.

Dark grey – signal peptide sequence. Brown – regions of intramolecular disulfide bonds. For simplicity only those regions coinciding with variants described were indicated. Black – bait region described to interact with proteases. Purple – thiol ester sequence region aiding in interaction with proteases. Light grey – alpha helical regions thought to mediate A2ML1 interaction with LRP1, facilitating receptor-mediated endocytosis.

**Table 1 Novel blood pressure trait associated loci. Variants with formal replication**

| Variant information |              |                     |       | Discovery              |                        | Replication |         |                        | Combined |         |                        |
|---------------------|--------------|---------------------|-------|------------------------|------------------------|-------------|---------|------------------------|----------|---------|------------------------|
| Locus               | rsID         | Chr:Pos (EA:EAF)    | Trait | $P_t$                  | $P_u$                  | N           | $\beta$ | $P$                    | N        | $\beta$ | $P$                    |
| <b>EUR</b>          |              |                     |       |                        |                        |             |         |                        |          |         |                        |
| <i>RNF207</i>       | rs709209     | 1:6.28 (A:0.655)    | PP    | $4.57 \times 10^{-6}$  | $1.60 \times 10^{-6}$  | 122,780     | 0.17    | $5.83 \times 10^{-4}$  | 284,683  | 0.20    | $9.62 \times 10^{-9}$  |
| <i>C5orf56</i>      | rs12521868   | 5:131.78 (T:0.373)  | DBP   | $1.59 \times 10^{-6}$  | $3.03 \times 10^{-7}$  | 122,795     | -0.18   | $2.29 \times 10^{-5}$  | 282,023  | -0.19   | $6.12 \times 10^{-11}$ |
| <i>PHACTR1</i>      | rs9349379    | 6:12.90 (A:0.566)   | SBP   | $2.11 \times 10^{-8}$  | $1.78 \times 10^{-7}$  | 122,809     | 0.24    | $4.06 \times 10^{-4}$  | 284,673  | 0.29    | $8.84 \times 10^{-10}$ |
| <i>COL21A1</i>      | rs200999181† | 6:55.94 (A:0.002)   | PP    | $3.08 \times 10^{-8}$  | $2.46 \times 10^{-7}$  | 121,487     | 2.70    | $1.90 \times 10^{-4}$  | 242,486  | 3.25    | $6.27 \times 10^{-10}$ |
| <i>ABO</i>          | rs687621     | 9:136.14 (A:0.615)  | DBP   | $8.80 \times 10^{-8}$  | $2.55 \times 10^{-7}$  | 122,798     | 0.16    | $1.96 \times 10^{-4}$  | 276,014  | 0.19    | $5.45 \times 10^{-10}$ |
| <i>ADO</i>          | rs10995311   | 10:64.56 (C:0.567)  | DBP   | $1.86 \times 10^{-6}$  | $1.14 \times 10^{-6}$  | 122,798     | 0.23    | $8.47 \times 10^{-8}$  | 266,456  | 0.21    | $1.12 \times 10^{-12}$ |
| <i>LMO1</i>         | rs110419     | 11:8.25 (A:0.48)    | DBP   | $9.41 \times 10^{-6}$  | $2.22 \times 10^{-5}$  | 122,798     | 0.16    | $1.81 \times 10^{-4}$  | 279,935  | 0.16    | $3.04 \times 10^{-8}$  |
| <i>OR5B12</i>       | rs11229457   | 11:58.21 (T:0.236)  | SBP   | $1.58 \times 10^{-6}$  | $4.62 \times 10^{-5}$  | 122,809     | -0.32   | $7.53 \times 10^{-5}$  | 284,680  | -0.31   | $2.70 \times 10^{-8}$  |
| <i>CERS5</i>        | rs7302981    | 12:50.54 (A:0.361)  | DBP   | $1.35 \times 10^{-13}$ | $4.60 \times 10^{-11}$ | 122,798     | 0.24    | $2.64 \times 10^{-8}$  | 284,718  | 0.25    | $1.38 \times 10^{-17}$ |
| <i>MYH6</i>         | rs452036     | 14:23.87 (A:0.327)  | PP    | $4.59 \times 10^{-11}$ | $2.80 \times 10^{-13}$ | 122,780     | -0.21   | $1.81 \times 10^{-5}$  | 284,672  | -0.28   | $2.96 \times 10^{-16}$ |
| <i>DPEP1</i>        | rs1126464    | 16:89.70 (C:0.256)  | DBP   | $1.19 \times 10^{-9}$  | $4.35 \times 10^{-11}$ | 118,677     | 0.24    | $1.68 \times 10^{-6}$  | 261,564  | 0.28    | $1.02 \times 10^{-15}$ |
| <i>TBX2</i>         | rs8068318†   | 17:59.48 (T:0.698)  | DBP   | $7.46 \times 10^{-13}$ | $5.71 \times 10^{-10}$ | 122,798     | 0.26    | $3.23 \times 10^{-8}$  | 281,978  | 0.26    | $1.95 \times 10^{-16}$ |
| <i>RGL3</i>         | rs167479     | 19:11.53 (T:0.486)  | DBP   | $2.22 \times 10^{-23}$ | $1.97 \times 10^{-22}$ | 122,797     | -0.29   | $3.01 \times 10^{-11}$ | 283,332  | -0.33   | $1.99 \times 10^{-31}$ |
| <i>PREX1</i>        | rs6095241    | 20:47.31 (A:0.452)  | DBP   | $5.65 \times 10^{-6}$  | $2.29 \times 10^{-5}$  | 122,798     | -0.18   | $2.56 \times 10^{-5}$  | 281,322  | -0.17   | $4.75 \times 10^{-9}$  |
| <b>ALL ancestry</b> |              |                     |       |                        |                        |             |         |                        |          |         |                        |
| <i>RBM47</i>        | rs35529250†  | 4:40.43 (T:0.01)    | SBP   | $6.56 \times 10^{-7}$  | $6.15 \times 10^{-6}$  | 148,878     | -1.43   | $5.02 \times 10^{-4}$  | 306,352  | -1.55   | $2.42 \times 10^{-8}$  |
| <i>OBFC1</i>        | rs4387287    | 10:105.68 (A:0.157) | SBP   | $2.23 \times 10^{-8}$  | $1.32 \times 10^{-7}$  | 147,791     | 0.28    | $3.37 \times 10^{-4}$  | 320,494  | 0.36    | $9.12 \times 10^{-10}$ |
| <i>RRAS</i>         | rs61760904†  | 19:50.14 (T:0.008)  | SBP   | $1.96 \times 10^{-6}$  | $1.90 \times 10^{-5}$  | 148,878     | 1.38    | $5.70 \times 10^{-4}$  | 322,664  | 1.50    | $8.45 \times 10^{-8}$  |

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SNV-BP associations are reported for the newly identified BP loci that replicated at  $P < 6.2 \times 10^{-4}$  (Bonferroni correction for the 81 variants selected for replication for a primary blood pressure trait; Methods). Loci are categorised into EUR and ALL ancestry based on the meta-analysis used to replicate the variants for the primary BP trait shown in columns labelled 'Trait'. In the columns that contains the discovery meta-analyses results,  $P_t$  represents the  $P$ -value for association of the variant with the transformed primary BP trait in the EUR\_SAS discovery meta-analyses (which was also used to select the variant for replication) and  $P_u$  represents the  $P$ -value for association with the untransformed primary BP trait in the ancestry in which the variant replicated. N,  $\beta$  and  $P$ , which denote the number of samples, estimated allelic effect and  $P$ -value for association with the primary BP trait, are provided for the untransformed primary BP trait in the replication data and also from the combined (discovery and replication) meta-analyses. NB: ALL ancestry corresponds to all ancestries in the combined (discovery + replication) meta-analyses

Locus – Gene or region containing the SNV, rsID – dbSNP rsID. Chr:Pos (EA:EAF) – Chromosome:NCBI Build 37 position in Mb (effect allele:effect allele frequency), Trait – primary blood pressure trait for which the variant was and also replicated,  $\beta$  – effect estimate, N:sample size, EUR – European.

† indicates it is a non-synonymous SNV (nsSNV) or is in linkage disequilibrium with a nsSNV ( $r^2 > 0.8$ ) that is predicted to be damaging

**Table 2 Novel blood pressure trait associated loci. Variants with GWS evidence of association in combined meta-analyses**

| Variant information |             |                     |                        | Discovery              |                       | Replication |         |                       | Combined |         |                        |
|---------------------|-------------|---------------------|------------------------|------------------------|-----------------------|-------------|---------|-----------------------|----------|---------|------------------------|
| Locus               | rsID        | Chr:Pos (EA:EAF)    | Trait                  | $P_t$                  | $P_u$                 | N           | $\beta$ | $P$                   | N        | $\beta$ | $P$                    |
| EUR                 |             |                     |                        |                        |                       |             |         |                       |          |         |                        |
| 2q36.3              | rs2972146   | 2:227.10 (T:0.652)  | DBP <sup>§</sup> (HTN) | $1.51 \times 10^{-9}$  | $2.47 \times 10^{-7}$ | 122,798     | 0.13    | $2.20 \times 10^{-3}$ | 275,610  | 0.17    | $8.40 \times 10^{-9}$  |
| ZBTB38              | rs16851397  | 3:141.13 (A:0.953)  | DBP <sup>§</sup> (SBP) | $6.87 \times 10^{-6}$  | $3.20 \times 10^{-5}$ | 122,798     | -0.38   | $1.20 \times 10^{-4}$ | 284,717  | -0.38   | $3.01 \times 10^{-8}$  |
| PRDM6               | rs1008058   | 5:122.44 (A:0.135)  | SBP                    | $5.09 \times 10^{-7}$  | $1.01 \times 10^{-8}$ | 43,109      | 0.46    | $3.61 \times 10^{-3}$ | 176,362  | 0.55    | $2.99 \times 10^{-10}$ |
| GPR20               | rs34591516  | 8:142.37 (T:0.055)  | SBP <sup>§</sup> (DBP) | $1.54 \times 10^{-6}$  | $1.01 \times 10^{-7}$ | 122,807     | 0.51    | $4.20 \times 10^{-4}$ | 282,009  | 0.64    | $6.10 \times 10^{-10}$ |
| HOXB7               | rs7406910   | 17:46.69 (T:0.118)  | SBP                    | $6.07 \times 10^{-10}$ | $2.74 \times 10^{-9}$ | 122,809     | -0.20   | $4.89 \times 10^{-2}$ | 284,690  | -0.46   | $3.80 \times 10^{-8}$  |
| AMH                 | rs10407022† | 19:2.25 (T:0.82)    | PP                     | $1.63 \times 10^{-7}$  | $1.73 \times 10^{-7}$ | 118,656     | -0.19   | $1.62 \times 10^{-3}$ | 252,525  | -0.26   | $5.94 \times 10^{-9}$  |
| ZNF101              | rs2304130   | 19:19.79 (A:0.914)  | DBP                    | $1.66 \times 10^{-8}$  | $1.92 \times 10^{-8}$ | 122,798     | -0.17   | $1.71 \times 10^{-2}$ | 284,705  | -0.29   | $1.53 \times 10^{-8}$  |
| PROCR               | rs867186    | 20:33.76 (A:0.873)  | DBP                    | $1.44 \times 10^{-6}$  | $4.15 \times 10^{-7}$ | 122,798     | 0.21    | $2.48 \times 10^{-3}$ | 284,722  | 0.26    | $1.19 \times 10^{-8}$  |
| RRP1B               | rs9306160   | 21:45.11 (T:0.374)  | DBP <sup>§</sup> (SBP) | $1.04 \times 10^{-8}$  | $1.90 \times 10^{-6}$ | 100,489     | -0.16   | $4.30 \times 10^{-4}$ | 249,817  | -0.18   | $6.80 \times 10^{-9}$  |
| TNRC6B              | rs470113    | 22:40.73 (A:0.804)  | PP                     | $1.48 \times 10^{-10}$ | $1.31 \times 10^{-9}$ | 122,780     | -0.14   | $1.37 \times 10^{-2}$ | 284,683  | -0.25   | $1.67 \times 10^{-9}$  |
| ALL ancestry        |             |                     |                        |                        |                       |             |         |                       |          |         |                        |
| 7q32.1              | rs4728142   | 7:128.57 (A:0.433)  | SBP                    | $8.10 \times 10^{-6}$  | $4.21 \times 10^{-6}$ | 150,542     | -0.21   | $8.62 \times 10^{-4}$ | 338,338  | -0.24   | $3.45 \times 10^{-8}$  |
| PRKAG1              | rs1126930†  | 12:49.40 (C:0.036)  | PP                     | $2.12 \times 10^{-6}$  | $4.62 \times 10^{-7}$ | 151,481     | 0.36    | $3.74 \times 10^{-3}$ | 314,894  | 0.50    | $3.34 \times 10^{-8}$  |
| SBNO1               | rs1060105   | 12:123.81 (T:0.209) | DBP                    | $6.66 \times 10^{-7}$  | $1.09 \times 10^{-6}$ | 150,532     | -0.15   | $2.67 \times 10^{-3}$ | 336,413  | -0.18   | $3.07 \times 10^{-8}$  |

SNV-BP associations are reported for the newly identified BP loci that showed genome-wide significant association ( $P < 5 \times 10^{-8}$ ) in the combined discovery and replication meta-analyses. In the columns that contain results from the discovery meta-analyses,  $P_t$  represents the  $P$ -value for association of the variant with the transformed *primary* BP trait in the EUR\_SAS discovery meta-analyses (used to select the variant for replication) and  $P_u$  represents the  $P$ -value for association with the untransformed BP trait in the ancestry in which the variant was validated. Loci are categorised into EUR and ALL ancestry based on the ancestry in which the variant showed association with a blood pressure trait at  $P < 5 \times 10^{-8}$ . N,  $\beta$  and  $P$ , which denote the number of samples, estimated allelic effect and  $P$ -value for association with the validated BP trait, are provided for the untransformed BP trait in the replication data and also from the combined (discovery and replication) meta-analyses. NB: ALL ancestry corresponds to all ancestries in the combined (discovery + replication) meta-analyses.

Locus – Gene or region containing the SNV, rsID - dbSNP rsID. Chr:Pos (EA:EAF) – Chromosome:NCBI Build 37 position in Mb (effect allele:effect allele frequency), Trait - blood pressure trait for which association is reported, EUR - European.

§ At four loci (2q36.3, ZBTB38, GPR20 and RRP1B) the primary trait used to select the variants for replication is given in parentheses because the variant associations were validated in the combined meta-analysis for the listed secondary trait. For these variants,  $P_t$  denotes the  $P$ -value for association with the primary trait, the other  $P$ -values provided are for the secondary trait.

† indicates it is a non-synonymous SNV (nsSNV) or is linkage disequilibrium with a nsSNV ( $r^2 > 0.8$ ) that is predicted to be damaging

206 **Table 3 Results of the genetic risk score analyses across CVD traits and risk factors.**

| Outcome                         | Units | N       | DBP (per 10mmHg increase) |                         | SBP (per 10mmHg increase) |                         | PP (per 10mmHg increase) |                         |
|---------------------------------|-------|---------|---------------------------|-------------------------|---------------------------|-------------------------|--------------------------|-------------------------|
|                                 |       |         | Effect [95% CI]           | P                       | Effect [95% CI]           | P                       | Effect [95% CI]          | P                       |
| CHD                             | OR    | 82,056  | 1.62 [ 1.28, 2.05]        | 5.99 x 10 <sup>-5</sup> | 1.39 [ 1.22, 1.59]        | 6.07 x 10 <sup>-7</sup> | 1.70 [ 1.34, 2.16]       | 1.20 x 10 <sup>-5</sup> |
| Ischemic stroke                 | OR    | 25,799  | 1.93 [ 1.47, 2.55]        | 2.81 x 10 <sup>-6</sup> | 1.57 [ 1.35, 1.84]        | 1.16 x 10 <sup>-8</sup> | 2.12 [ 1.58, 2.84]       | 5.35 x 10 <sup>-7</sup> |
| Cardioembolic stroke            | OR    | 16,113  | 1.43 [ 0.86, 2.39]        | 0.1683                  | 1.33 [ 0.99, 1.80]        | 0.0584                  | 1.73 [ 1.00, 3.02]       | 0.0518                  |
| Large vessel stroke             | OR    | 13,903  | 2.26 [ 1.25, 4.08]        | 0.0068                  | 1.85 [ 1.32, 2.59]        | 3.61 x 10 <sup>-4</sup> | 3.05 [ 1.64, 5.68]       | 4.37 x 10 <sup>-4</sup> |
| Small vessel stroke             | OR    | 15,617  | 1.96 [ 1.13, 3.41]        | 0.0168                  | 1.56 [ 1.13, 2.16]        | 0.0064                  | 1.98 [ 1.09, 3.61]       | 0.0248                  |
| Heart failure                   | OR    | 13,282  | 1.48 [ 1.02, 2.17]        | 0.0409                  | 1.25 [ 1.00, 1.57]        | 0.0512                  | 1.33 [ 0.88, 2.02]       | 0.1757                  |
| Left ventricular mass           | g     | 11,273  | 9.57 [ 3.98,15.17]        | 8.02 x 10 <sup>-4</sup> | 5.13 [ 1.77, 8.48]        | 0.0027                  | 5.97 [-0.38,12.31]       | 0.0653                  |
| Left ventricular wall thickness | cm    | 11,311  | 0.10 [ 0.06, 0.13]        | 1.88 x 10 <sup>-8</sup> | 0.05 [ 0.03, 0.07]        | 5.52 x 10 <sup>-6</sup> | 0.05 [ 0.01, 0.09]       | 0.0187                  |
| HDL                             | mg/dl | 80,395  | 0.25 [-1.00, 1.51]        | 0.6930                  | 0.21 [-0.50, 0.92]        | 0.5622                  | 0.47 [-0.79, 1.73]       | 0.4668                  |
| LDL                             | mg/dl | 77,021  | -1.57 [-5.20, 2.06]       | 0.3972                  | 0.07 [-2.03, 2.16]        | 0.9498                  | 1.87 [-1.86, 5.59]       | 0.3255                  |
| Total cholesterol               | mg/dl | 80,455  | -1.34 [-5.90, 3.22]       | 0.5639                  | 0.70 [-1.93, 3.32]        | 0.6029                  | 3.68 [-0.97, 8.33]       | 0.1209                  |
| Triglycerides                   | mg/dl | 77,779  | 0.02 [-0.03, 0.08]        | 0.3859                  | 0.02 [-0.01, 0.05]        | 0.2697                  | 0.03 [-0.03, 0.08]       | 0.3025                  |
| BMI                             | INVT  | 526,508 | -0.10 [-0.18,-0.01]       | 0.0342                  | -0.07 [-0.13,-0.02]       | 0.0058                  | -0.12 [-0.23,-0.02]      | 0.0165                  |
| WHRadjBMI                       | INVT  | 344,369 | 0.03 [-0.04, 0.11]        | 0.4025                  | 0.03 [-0.02, 0.08]        | 0.2170                  | 0.06 [-0.03, 0.15]       | 0.1885                  |
| Height                          | INVT  | 458,927 | 0.02 [-0.15, 0.18]        | 0.8592                  | -0.04 [-0.15, 0.06]       | 0.4170                  | -0.18 [-0.37, 0.01]      | 0.0683                  |
| eGFR                            | INVT  | 51,039  | -0.02 [-0.15, 0.11]       | 0.7810                  | -0.03 [-0.10, 0.04]       | 0.4080                  | -0.07 [-0.20, 0.06]      | 0.2741                  |

207 CHD, coronary heart disease; HDL, high density lipoprotein; LDL, low density lipoprotein; eGFR, estimated glomerular filtration rate; DBP, diastolic blood pressure; SBP systolic blood pressure; PP, pulse pressure; OR, odds ratio; g, grams; INVT, inverse normally  
208 transformed (hence no units); N, sample size; P, P-value of association of BP with the trait listed; CI, confidence interval. Results are considered significant if  $P < 0.0038$ , which corresponds to a Bonferroni correction for 13 phenotypes tested.



## 211 **Online Methods**

### 212 **Overview of discovery studies**

213 The cohorts contributing to the discovery meta-analyses comprise studies from three consortia (CHD  
214 Exome+, ExomeBP, and GoT2D/T2D-GENES) with a total number of 192,763 unique samples. All  
215 participants provided written informed consent and the studies were approved by their local Research Ethics  
216 Committees and/or Institutional Review Boards.

217 The CHD Exome+ consortium comprised 77,385 samples: eight studies (49,898 samples) of European  
218 (EUR) ancestry, two studies (27,487 samples) of South Asian (SAS) ancestry (Supplementary Table 1). The  
219 ExomeBP consortium included 25 studies (75,620 samples) of EUR ancestry (Supplementary Table 1). The  
220 GoT2D consortium comprised 14 studies (39,758 samples) of Northern EUR ancestry from Denmark,  
221 Finland, and Sweden (Supplementary Table 1). The participating studies and their characteristics including  
222 BP phenotypes are detailed in Supplementary Tables 1 and 2. Note, any studies contributing to multiple  
223 consortia were only included once in all meta-analyses.

### 224 **Phenotypes**

225 Four blood pressure (BP) traits were analysed: systolic blood pressure (SBP), diastolic blood pressure  
226 (DBP), pulse pressure (PP) and hypertension (HTN). For individuals known to be taking BP lowering  
227 medication, 15/10 mmHg was added to the raw SBP/DBP values, respectively, to obtain medication-  
228 adjusted SBP/DBP values<sup>45</sup>. PP was defined as SBP minus DBP, post-adjustment. For HTN, individuals  
229 were classified as hypertensive cases if they satisfied at least one of: (i)  $SBP \geq 140$  mmHg, (ii)  $DBP \geq 90$   
230 mmHg, (iii) taking anti-hypertensive or BP lowering medication. All other individuals were included as  
231 controls. The four BP traits were correlated (SBP:DBP correlations were between 0.6 and 0.8, and SBP:PP  
232 correlations were  $\sim 0.8$ ). However, they measure partly distinct physiological features including, cardiac  
233 output, vascular resistance, and arterial stiffness, all measures for determining a cardiovascular risk profile.  
234 Therefore the genetic architecture of the individual phenotypes are of interest, and a multi-phenotype  
235 mapping approach was not adopted.

236

## 238 **Genotyping**

239 All samples were genotyped using one of the Illumina HumanExome Beadchip arrays (Supplementary Table  
240 3). An Exome chip quality control Standard Operating Procedure (SOP) developed by Anubha Mahajan,  
241 Neil Robertson and Will Rayner at the Wellcome Trust Centre for Human Genetics, University of Oxford  
242 was used by most studies for genotype calling and QC<sup>46</sup> (Supplementary Table 3). All genotypes were  
243 aligned to the plus strand of the human genome reference sequence (Build37) prior to any analyses and any  
244 unresolved mappings were removed. Genotype cluster plots were reviewed for all the novel rare variants  
245 (both lead and secondary signals) and for rare variants that contributed to the gene-based testing.

## 246 **Meta-analyses**

247 Meta-analyses were performed using METAL<sup>47</sup>, for both discovery and replication analyses, using inverse  
248 variance weighted fixed effect meta-analysis for the continuous traits (SBP, DBP and PP) and sample size  
249 weighted meta-analysis for the binary trait (HTN).

## 250 **Discovery SNV analyses**

251 Analyses of both untransformed and inverse normal transformed SBP, DBP and PP were conducted within  
252 each contributing study. The analyses of the transformed traits were performed in order to minimise  
253 sensitivity to deviations from normality in the analysis of rare variants and for discovery of new SNV-BP  
254 associations. The residuals from the null model obtained after regressing the medication-adjusted trait on the  
255 covariates (age, age<sup>2</sup>, sex, BMI, and disease status for CHD) within a linear regression model, were ranked  
256 and inverse normalised. These normalised residuals were used to test trait-SNV associations. All SNVs that  
257 passed QC were analysed for association, without any further filtering by MAF, but a minor allele count of  
258 10 was used for the analysis of HTN. An additive allelic effects model was assumed.

259 Two meta-analyses were performed for each trait, one with EUR and SAS ancestries combined (EUR\_SAS)  
260 and another for EUR ancestry alone. Contributing studies used principal components (PCs) to adjust for  
261 population stratification. Consequently minimal inflation in the association test statistics,  $\lambda$ , was observed (

262  $\lambda = 1.07$  for SBP, 1.10 for DBP, 1.04 for PP and  $<1$  for HTN in the transformed discovery meta-analysis in  
263 EUR\_SAS;  $\lambda = 1.06$  for SBP, 1.09 for DBP, 1.05 for PP and  $<1$  for HTN in the transformed discovery  
264 meta-analysis in EUR; Supplementary Figure 6). The meta-analyses were performed independently in two  
265 centres and results were found to be concordant between centres. Given the studies contributing to the  
266 discovery analyses were ascertained on CHD or T2D, we tested potential systematic bias in calculated effect  
267 estimates amongst these studies. No evidence of bias in the overall effect estimates was obtained.

268 The results for the transformed traits were taken forward and used to select candidate SNVs for replication.  
269 Results ( $P$ -values) from the transformed and untransformed analyses were strongly correlated ( $r^2 > 0.9$ ).

## 270 **Replication SNV analyses**

271 SNVs associated with any of the transformed traits (SBP, DBP, PP) or HTN were annotated using the  
272 Illumina SNV annotation file, humanexome-12v1\_a\_gene\_annotation.txt, independently across two centres.  
273 Given the difference in power to detect common versus low frequency and rare variant associations, two  
274 different significance thresholds were chosen for SNV selection. For SNVs with  $MAF \geq 0.05$ ,  $P \leq 1 \times 10^{-5}$  was  
275 selected, while,  $P \leq 1 \times 10^{-4}$  was used for SNVs with  $MAF < 0.05$ . By choosing a significance threshold of  
276  $P < 1 \times 10^{-4}$  we maximized the opportunity to follow-up rare variants (making the assumption that any true  
277 signals at this threshold could replicate at Bonferroni adjusted significance,  $P \leq 6.17 \times 10^{-4}$ , assuming  $\alpha = 0.05$   
278 for 81 SNVs). All previously published BP associated SNVs and any variants in LD with them ( $r^2 > 0.2$ ),  
279 were removed from the list of associated SNVs as we aimed to replicate new findings only. SNVs for which  
280 only one study contributed to the association result or showed evidence of heterogeneity ( $P_{het} < 0.0001$ ) were  
281 removed from the list as they were likely to be an artefact. Where SNVs were associated with multiple traits,  
282 to minimise the number of tests performed, only the trait with the smallest  $P$ -value was selected as the  
283 primary trait in which replication was sought. Where multiple SNVs fitted these selection criteria for a  
284 single region, only the SNV with the smallest  $P$ -value was selected. In total, 81 SNVs were selected for  
285 validation in independent samples. These 81 SNVs had concordant association results for both transformed  
286 and non-transformed traits. Eighty SNVs were selected from EUR\_SAS results (with consistent support in  
287 EUR), and one SNV from EUR results only. In the next step, we looked up the 81 SNV-BP associations  
288 using data from a separate consortium, the CHARGE+ exome chip blood pressure consortium (who had



289 analysed untransformed SBP, DBP, PP and HTN), and UHP and Lolipop (ExomeBP consortium;  
290 Supplementary Tables 2 and 3). The analysed residuals from CHARGE+ were approximately normally  
291 distributed in their largest studies (Supplementary Figure 7).

292 Two meta-analyses of the replication datasets were performed: one of EUR samples, and a second of EUR,  
293 African American, Hispanics and SAS ancestries (“ALL”). Replication was confirmed if  $P$  (1-tailed)  $<$   
294  $0.05/81=6.17 \times 10^{-4}$  and the effect (beta) was in the direction observed in discovery meta-analyses for the  
295 selected trait. A combined meta-analysis was performed of discovery (untransformed results as only  
296 untransformed data was available from CHARGE+ exome chip blood pressure consortium) and replication  
297 results across the four traits to assess the overall support for each locus. For the combined meta-analyses, a  
298 GWS threshold of,  $P \leq 5 \times 10^{-8}$ , was used to declare a SNV as novel rather than a less stringent experiment  
299 wide threshold, as GWS is used to declare significance in GWAS and we wish to minimise the possibility of  
300 false positive associations. (Note that GWS is equivalent to an exome-wide threshold of  $P \leq 2 \times 10^{-7}$  adjusted  
301 for four traits).

302

303 Note: all validated BP-associated variants were associated at  $P < 10^{-5}$  in the discovery dataset (for the primary  
304 trait). Hence, we could have used the same inclusion criteria for both common and rare SNVs. Therefore the  
305 optimal threshold to choose for future experiments may need further consideration.

### 306 **Conditional analyses and gene-based tests**

307 The RAREMETALWORKER (RMW) tool<sup>15</sup> (version 4.13.3) that does not require individual level data to  
308 perform conditional analyses and gene-based tests was used for conditional analyses. All studies that  
309 contributed to the SNV discovery analyses were re-contacted and asked to run RMW. Only FENLAND,  
310 GoDARTS, HELIC-MANOLIS, UKHLS and EPIC-InterAct were unable to run RMW, while two new  
311 studies were included, INCIPE and NFBC1966 (Supplementary Table 1 and 2). In total, 43 studies (147,402  
312 samples) were included in the EUR analyses and 45 studies (173,329 samples) in the EUR\_SAS analyses  
313 (Supplementary Tables 2 and 3). Comparison of discovery and RMW study level results were made  
314 (Supplementary Information).

315 For each novel locus, the genomic coordinates and size of the region were defined according to  
316 recombination rates (Supplementary Table 9) around the lead variant. For known loci, a 1 Mb window was  
317 used (Supplementary Table 14). Conditional analyses were performed across each region, in both EUR and  
318 EUR\_SAS samples, for the transformed phenotype corresponding to the validated BP trait for novel loci and  
319 the published BP trait for known loci.

320 Gene based tests were performed in both the EUR and EUR\_SAS datasets using the Sequence Kernel  
321 Association Test (SKAT)<sup>16</sup> method implemented in RMW as it allows for the SNVs to have different  
322 directions and magnitudes of effect. Burden tests were also performed but are not presented as only SKAT  
323 provided significant results. The variants in the gene-based tests using SKAT were weighted using the  
324 default settings, *i.e.* a beta distribution density function to up-weight rare variants,  $\text{Beta}(\text{MAF}_j, 1, 25)$  where  
325  $\text{MAF}_j$  represents the pooled MAF for variant  $j$  across all studies. Analyses were restricted to coding SNVs  
326 with  $\text{MAF} < 5\%$  and  $< 1\%$ . Genes were deemed to be associated if  $P < 2.8 \times 10^{-6}$  (Bonferroni adjusted for  
327 17,996 genes). To confirm the gene associations were not attributable to a solitary SNV, a gene-based test  
328 conditional on the most associated SNV was performed ( $P_{\text{conditional}} < 0.001$ ). The QC of all SNVs  
329 contributing to the gene based tests including the number of samples and studies were checked prior to  
330 claiming association. We sought replication of associated genes in the CHARGE+ exome chip blood  
331 pressure consortium.

332

### 333 **Pathway analyses with MAGENTA**

334 We tested seven databases in MAGENTA<sup>37</sup> (BioCarta, Kyoto Encyclopedia of Genes and Genomes,  
335 Ingenuity, Panther, Panther Biological Processes, Panther Molecular Functions and Reactome) for  
336 overrepresentation of the SNV discovery results from both EUR and EUR\_SAS ancestries. Each of the four  
337 BP phenotypes were tested. Pathways exhibiting  $P < 0.01$  and  $\text{FDR} < 5\%$  were considered statistically  
338 significant.

### 339 **GeneGo MetaCore Network analyses**

A set of BP genes based on previously published studies and our current results (locus defined as  $r^2 > 0.4$  and 500kb on either side of the lead SNV; Supplementary Table 19) were tested for enrichment using the THOMSON REUTERS MetaCore™ Single Experiment Analysis workflow tool. The data were mapped onto selected MetaCore ontology databases: pathway maps, process networks, GO processes and diseases / biomarkers, for which functional information is derived from experimental literature. Outputs were sorted based on  $P$ - and FDR-values. A gene set was considered enriched for a particular process if  $P < 0.05$  and  $FDR < 5\%$ .

### Genetic Risk Score

To assess the effect of BP on CHD, ischemic stroke (and subtypes: large vessel, small vessel and cardioembolic stroke) left ventricular mass, left ventricular wall thickness, heart failure, HDL-c, LDL-c, total cholesterol, triglycerides and eGFR, we performed a weighted generalized linear regression of the genetic associations with each outcome variable on the genetic associations with BP. When genetic variants are uncorrelated, the estimates from such a weighted linear regression analysis using summarized data, and a genetic risk score analysis using individual-level data, are equal<sup>48</sup>. We refer to the analysis as a genetic risk score (also known as a polygenic risk score) analysis as this is likely to be more familiar to applied readers. As some of the genetic variants in our analysis are correlated, a generalized weighted linear regression model is fitted that accounts for the correlations between variants, as follows: If  $\beta_X$  are the genetic associations (beta-coefficients) with the risk factor (here, BP) and  $\beta_Y$  are the genetic associations with the outcome, then the causal estimate from a weighted generalized linear regression is  $(\beta_X^T \Omega^{-1} \beta_X)^{-1} \beta_X^T \Omega^{-1} \beta_Y$ , with standard error,

$$\hat{\sigma} \sqrt{(\beta_X^T \Omega^{-1} \beta_X)^{-1}},$$

where  $^T$  is a matrix transpose,  $\hat{\sigma}$  is the estimate of the residual standard error from the regression model, and the weighting matrix  $\Omega$  has terms

$$\Omega_{j_1 j_2} = \sigma_{Y j_1} \sigma_{Y j_2} \rho_{j_1 j_2}$$

, where  $\sigma_{Y j}$  is the standard error of the genetic association with the outcome for the  $j$ th SNV, and  $\rho_{j_1 j_2}$  is the correlation between the  $j_1$ th and  $j_2$ th SNVs. The presence of the estimated residual standard error allows for

heterogeneity between the causal estimates from the individual SNVs as overdispersion in the regression model (in the case of underdispersion, the residual standard error estimate is set to unity). This is equivalent to combining the causal estimates from each SNV using a multiplicative random-effects model<sup>49</sup>.

For each of SBP, DBP and PP, the score was created using both the novel and known BP SNVs or a close proxy ( $r^2 > 0.8$ ). Both the sentinel SNV association and any secondary SNV associations that remained after adjusting for the sentinel SNV were included in the genetic risk score. For the 30 validated novel SNV-BP associations,  $\beta$ s were taken from the independent replication analyses (Table 1 and 2) to weight the SNV in the genetic risk score. For the secondary SNVs from the seven novel loci and five known loci,  $\beta$ s were taken from the discovery analyses (Supplementary Tables 10 and 15). For the 82 known SNVs, 43 were either genotyped or had proxies on the Exome chip and the  $\beta$ s were taken from discovery results (Supplementary Table 13), the remaining  $\beta$ s were taken from published effect estimates. This strategy for selecting betas for use in the GRS was taken to minimize the influence of winner's curse. The associations between the BP variants with CHD, HDL-c, LDL-c, total cholesterol, log(triglycerides) and log(eGFR) were obtained using the CHD Exome+ Consortium studies, the associations with BMI, waist-hip ratio adjusted BMI and height from the GIANT consortium (unpublished data), ischemic stroke from METASTROKE<sup>25</sup>, and left ventricular mass, left ventricular wall thickness and heart failure from EchoGen<sup>27</sup> and CHARGE-HF<sup>26</sup>. A causal interpretation of the association of GRS with the outcome as the effect of BP on the outcome assumes that the effects of genetic variants on the outcome are mediated via blood pressure and not via alternate causal pathways, for example via LV thickness. There are also limitations of the Mendelian randomization approach in distinguishing between the causal effects of different measures of blood pressure, due to the paucity of genetic variants associated with only one measure of blood pressure.

## **eQTL analyses**

The MuTHER dataset contains gene expression data from 850 UK twins for 23,596 probes and 2,029,988 (HapMap 2 imputed) SNVs. All cis-associated SNVs with FDR < 1%, within each of the 30 novel regions (IMPUTE info score > 0.8) were extracted from the MuTHER project dataset for, LCL (n=777), adipose

(n=776) and skin (n=667)<sup>50</sup>. The pilot phase of the GTEx Project (dbGaP Accession phs000424.v3.p1) provides expression data from up to 156 individuals for 52,576 genes and 6,820,472 genotyped SNVs (imputed to 1000 Genomes project, MAF $\geq$ 5%)<sup>51</sup>. The eQTL analysis was focused on subcutaneous adipose tissue (n=94), tibial artery (n=112), heart (left ventricle) (n=83), lung (n=119), skeletal muscle (n=138), tibial nerve (n=88), skin (sun exposed, lower leg) (n=96), thyroid (n=105) and whole blood (n=156) which have >80 samples and genes expressed at least 0.1 RPKM in 10 or more individuals in a given tissue. All transcripts with a transcription start site (TSS) within one of the 30 new BP loci and for which there was a cis-associated SNV (IMPUTE info score >0.4) within 1Mb of the TSS at FDR<5%, were identified. Kidney was not evaluated because the sample size was too small (n=8). From each resource, we report eQTL signals, which reach the resource-specific thresholds for significance described above, for SNVs that are in LD ( $r^2$ >0.8) with our sentinel SNV.

For identified eQTLs, we tested whether they colocalised with the BP associated SNV<sup>52</sup>. Colocalisation analyses were considered to be significant if the posterior probability of colocalisation was greater than 0.95.

## Annotation of variants

*In silico* prediction of the functional effect of associated variants was based on the annotation from dbSNP, the Ensembl Variant Effect Predictor tool and the Exome Variant Server, NHLBI GO Exome Sequencing Project (ESP), Seattle, WA.

## Trait variance explained

The percentage trait variance explained for SBP, DBP, PP was assessed with 5,861 individuals with complete information for all phenotypes and covariates from the population-based cohort, 1958BC.

Two genetic models were investigated: one containing the 43 previously known BP associated SNVs covered on the Exome chip; the other additionally including the 30 novel lead SNVs and 9 conditionally independent SNVs from both novel and known loci. These nine conditionally independent SNVs were taken from the EUR results, as 1958BC is EUR. They included four from novel loci (*PREX1*, *COL21A1*, *PRKAG1* and *MYH6* (there was only 1 in EUR); Supplementary Table 10) and five from known loci (*ST7L-CAPZA1-MOV10*, *FIGN-GRB14*, *ENPEP*, *TBX5-TBX3* and *HOXC4*; Supplementary Table 15).

The residual trait was obtained by adjusting each of the BP traits in a regression model with sex and BMI variables (not age or age<sup>2</sup> as all 1958BC individuals were aged 44 years). The residual trait was regressed on all SNVs within the corresponding model and adjusted for the first ten PCs. The R<sup>2</sup> calculated from this regression model was used as the percentage trait variance explained.

## Monogenic Enrichment analyses

To determine if sub-significant signals of association were present in a set of genes associated with monogenic forms of disease, we performed an enrichment analysis of the discovery single variant meta-analyses association results for all four traits, both for EUR and EUR\_SAS datasets.

The monogenic gene set included: *WNK1*, *WNK4*, *KLHL3*, *CUL3*, *PPARG*, *NR3C2*, *CYP11B1*, *CYP11B2*, *CYP17A1*, *HSD11B2*, *SCNN1A*, *SCNN1B*, *SCNN1G*, *CLCNKB*, *KCNJ1*, *SLC12A1*, *SLC12A3*<sup>3</sup>. The association results of coding SNVs in these genes were extracted and the number of tests with  $P < 0.001$  observed. In order to determine how often such an observation would be observed by chance, we constructed 1,000 matched gene sets. The matching criteria for each monogenic gene was the intersection of all genes in the same exon length quintile and all genes in the same coding variant count decile. Within the matched sets, the number of variants with  $P < 0.001$  was observed. The empirical  $P$ -value was calculated as the fraction of matched sets with an equal or larger number of variants less than 0.001.

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